

Factors associated with colorectal cancer occurrence after colonoscopy that did not diagnose colorectal cancer

Cheung, Danny; Evison, Felicity; Patel, Prashant; Trudgill, Nigel

DOI:

[10.1016/j.gie.2016.01.047](https://doi.org/10.1016/j.gie.2016.01.047)

License:

None: All rights reserved

Document Version

Peer reviewed version

Citation for published version (Harvard):

Cheung, D, Evison, F, Patel, P & Trudgill, N 2016, 'Factors associated with colorectal cancer occurrence after colonoscopy that did not diagnose colorectal cancer', *Gastrointestinal Endoscopy*, vol. 84, no. 2, pp. 287-295.e1. <https://doi.org/10.1016/j.gie.2016.01.047>

[Link to publication on Research at Birmingham portal](#)

General rights

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

- Users may freely distribute the URL that is used to identify this publication.
- Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.
- User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?)
- Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

Manuscript Number: GIE-D-15-01290R2

Title: Factors associated with colorectal cancer occurrence after colonoscopy that did not diagnose colorectal cancer

Article Type: Original Article

Keywords: Colonoscopy; colorectal cancer

Corresponding Author: Dr. Danny W F Cheung, MBBS

Corresponding Author's Institution: Sandwell General Hospital

First Author: Danny W F Cheung, MBBS

Order of Authors: Danny W F Cheung, MBBS; Felicity Evison; Prashant Patel; Nigel Trudgill

Manuscript Region of Origin: UNITED KINGDOM (Scotland, Northern Ireland, Wales, Isle of Man, Channel Islands)

Abstract: Background and Aims: Up to 6% of colorectal cancers (CRC) are diagnosed within 5 years of a colonoscopy that did not diagnose CRC (post-colonoscopy colorectal cancer, PCCRC). PCCRC and associated risk factors were examined within a national hospital episode database.

Methods: A retrospective case-control study of all adult colonoscopies recorded in Hospital Episode Statistics (HES) between 2003-2009 in England. PCCRC cases underwent colonoscopy 6-60 months before diagnosis; controls had not undergone colonoscopy 6-60 months before diagnosis. Multivariate logistic regression analysis examined associations with PCCRC.

Results: 1,439,684 colonoscopies were analysed, including 67,202 CRC and 8147 (12.1%) PCCRC cases. Multivariate analysis revealed that female gender (odds ratio 1.13 (95% CI 1.08-1.19), $p<0.001$), older age (70-74 years) (1.09 (1.00-1.18), $p=0.039$), increased co-morbidity (Charlson index 5+) (1.16 (1.05-1.28), $p<0.003$) and right sided CRC (1.17 (1.11-1.23), $p<0.0001$) were associated with PCCRC. Emergency colonoscopy (0.54 (0.59-0.69), $p<0.0001$) was negatively associated with PCCRC. More PCCRC subjects developed metastases within 12 months and less underwent surgery (0.33 (0.32-0.35), $p<0.0001$) or chemotherapy (0.66 (0.62-0.69), $p<0.0001$). PCCRC rates varied twofold between providers, and was associated with medium volume providers compared with high volume (1.13 (1.01-1.27), $p=0.035$). The PCCRC rate fell from 13.8% in 2003 to 11.9% in 2009.

Conclusions: PCCRC occurred in 12.1% of CRC patients between 2003 and 2009. PCCRC was associated with female gender, older age, increased co-morbidity, right sided CRC, elective procedures and colonoscopy volume. PCCRC was associated with worse outcomes.

**Factors associated with colorectal cancer occurrence after colonoscopy that did not
diagnose colorectal cancer.**

¹Danny Cheung MBBS, ²Felicity Evison MSc, ³Prashant Patel MBBS PhD, ¹Nigel Trudgill MBBS MD

¹Sandwell General Hospital, Lyndon, West Bromwich, United Kingdom

²Health Informatics Department, Queen Elizabeth Hospital, Birmingham

³School of Cancer Sciences, University of Birmingham, Birmingham

Corresponding author:

Dr Nigel J Trudgill

Sandwell General Hospital

Lyndon

West Bromwich B71 4HJ

United Kingdom

Tel 0121 5073080

Fax 0121 5073265

Email nigel.trudgill@nhs.net

Word count: 3561

**Factors associated with colorectal cancer occurrence after colonoscopy that did not
diagnose colorectal cancer**

¹Danny Cheung, ²Felicity Evison, ³Prashant Patel, ¹Nigel Trudgill

¹Department of Gastroenterology, Sandwell General Hospital, Lyndon, West Bromwich

²Health Informatics Department, Queen Elizabeth Hospital, Birmingham

³School of Cancer Sciences, University of Birmingham, Birmingham

Corresponding author:

Dr NJ Trudgill

Sandwell General Hospital

Lyndon

West Bromwich

B71 4HJ

Tel 0121 5073080

Fax 0121 5073265

Email nigel.trudgill@nhs.net

Word count: 3561

Abstract

Background and Aims: Up to 6% of colorectal cancers (CRCs) are diagnosed within 5 years of a colonoscopy that did not diagnose CRC (post-colonoscopy colorectal cancer, PCCRC). PCCRC and associated risk factors were examined within a national hospital episode database.

Methods: A retrospective case-control study of all adult colonoscopies recorded in Hospital Episode Statistics (HES) between 2003 and 2009 in England. PCCRC cases underwent colonoscopy 6 to 60 months before diagnosis; controls had not undergone colonoscopy 6 to 60 months before diagnosis. Multivariate logistic regression analysis examined associations with PCCRC.

Results: A total of 1,439,684 colonoscopies were analyzed, including 67,202 CRC and 8147 (12.1%) PCCRC cases. Multivariate analysis revealed that female gender (odds ratio [OR], 1.13; 95% CI, 1.08-1.19), older age (70-74 years) (OR, 1.09; 95% CI, 1.00-1.18), $p=0.039$), increased co-morbidity (Charlson index 5+) (OR, 1.16; 95% CI, 1.05-1.28), $p<0.003$) and right-sided CRC (OR, 1.17; 95% CI, 1.11-1.23), $p<0.0001$) were associated with PCCRC. Emergency colonoscopy (OR, 0.54; 95% CI, 0.59-0.69), $p<0.0001$) was negatively associated with PCCRC. More PCCRC subjects developed metastases within 12 months and fewer underwent surgery (OR, 0.33; 95% CI, 0.32-0.35), $p<0.0001$) or chemotherapy (OR, 0.66; 95% CI, 0.62-0.69), $p<0.0001$). PCCRC rates varied twofold between providers and was associated with medium volume providers compared with high volume (OR, 1.13; 95% CI, 1.01-1.27), $p=0.035$). The PCCRC rate fell from 13.8% in 2003 to 11.9% in 2009.

Conclusions: PCCRC occurred in 12.1% of CRC patients between 2003 and 2009. PCCRC was associated with female gender, older age, increased co-morbidity, right-sided CRC, elective procedures, and colonoscopy volume. PCCRC was associated with worse outcomes.

Introduction

Colonoscopy is the criterion standard for diagnosing, screening and surveillance for CRC. In England, the setting of national standards for colonoscopy and accreditation of endoscopy units has resulted in an improvement in auditable colonoscopy standards over the last decade.[1] The same period has also coincided with an increase in 5-year survival after CRC diagnosis from 47.8% to 53.6%.[2] However, 2.6% to 6.0% of CRC patients have previously been reported to be diagnosed within 5 years of a colonoscopy that did not detect cancer. These events are termed post-colonoscopy colorectal cancer (PCCRC).[3, 4, 5] It has been proposed that PCCRC may have a different cell biology from other CRC with more aggressive and rapidly growing tumors.[6, 7] However, 2 recently published North American studies concluded that this did not apply to the majority of PCCRC, with around two-thirds of PCCRC a result of missed lesions or incomplete polypectomy.[4, 8]

Given the improvements in colonoscopy over the past decade in England, we have examined the impact on PCCRC in a national hospital episode database and associated risk factors for these events.

Methods

Data sources

Hospital Episode Statistics (HES) is an administrative database that records information on all elective and emergency care episodes in National Health Service (NHS) hospitals in England.[9] Each care episode record includes demographic, admission, diagnoses and procedures data. Diagnoses are coded using International Classification of Diseases version 10 (ICD-10) and procedures are coded using Office of Population Censuses and Surveys Classification of Interventions and Procedures 4th revision (OPCS-4). HES is linked to Office for National Statistics (ONS) mortality records, which include date of death and causes of death recorded on death certificates. The NHS provides comprehensive healthcare coverage for the UK population, with the vast majority of colonoscopies performed in the UK by a NHS provider.[1]

Subject definitions

All subjects over the age of 18 years undergoing colonoscopy between April 2003 and March 2009 were identified from HES. Colonoscopy and CRC were defined by OPCS-4 (*appendix 1*) and ICD-10 codes (*appendix 2*) respectively. Subjects with a CRC diagnosis before the first episode of colonoscopy and subjects with a diagnosis of inflammatory bowel disease (IBD) were excluded from the analysis to avoid confounding through surveillance.

Recording of a CRC diagnosis in HES records may be delayed by a few weeks from the date of the diagnostic colonoscopy code.[10, 11] For the purpose of this study, the diagnosis date was therefore defined as the first colonoscopy code during the 6 months before the first CRC coding episode in HES or mortality records[10, 12], or the first CRC episode for those subjects who did not have a colonoscopy during this 6-month period due to being diagnosed through an alternative method, eg, barium enema, CT colonography or flexible sigmoidoscopy. Subjects undergoing colonoscopy 6 to 60 months before subsequent CRC diagnosis were identified as PCCRC cases. These cases were further classified as PCCRC 6 to 12 months (colonoscopy 6 to 12 months before CRC diagnosis); PCCRC 12 to 36 months (colonoscopy 12 to 36 months before CRC diagnosis) and PCCRC 36 to 60 months (colonoscopy 36 to 60 months before CRC diagnosis). For patients who had more than one colonoscopy 6 to 60 months before CRC diagnosis, data from the most recent colonoscopy was used for analysis. Controls were subjects who had not undergone colonoscopy in the period 6 to 60 months before CRC diagnosis. Colonoscopies from 2003 to 2009 were studied to ensure all subjects had at least 5 years of follow-up within HES. The PCCRC rate was calculated from the number of PCCRC subjects divided by the sum of PCCRC subjects and controls.[13]

Validation of colonoscopy and colorectal cancer populations

To assess the validity of the HES colonoscopy population, the number of colonoscopies between 2007 and 2010 at University Hospital Birmingham (UHB) was extracted from endoscopy records (Unisoft Medical Systems, Enfield, Middlesex, UK) and compared with the number of colonoscopies recorded in HES for UHB. To assess the validity of a CRC

diagnosis in HES using the study methodology, the number of HES CRC cases was compared with the number of CRC cases diagnosed in England from the National Cancer Intelligence Network (NCIN)[14] from 2002 to 2011. Finally, the rate of surgery in the HES CRC population was compared with rate of surgery in the National Bowel Cancer Audit between 2008 and 2011.[15, 16, 17]

Study variables

Subject demographics

Study variables were extracted from coding at the time of PCCRC colonoscopy in cases and diagnostic colonoscopy or first CRC episode in controls. Ethnicity was identified from HES demographic fields and grouped into White or White British, Asian or Asian British, Black or Black British, Chinese, Mixed and other ethnic groups.

Co-morbidity

The Charlson co-morbidity index was calculated using ICD-10 codes for secondary diagnoses, excluding metastatic disease, and divided into 3 categories: 0 (no co-morbidity), 1 to 4 (low co-morbidity) and 5 or greater (high co-morbidity).[18]

Socio-economic status

Deprivation was assessed using the Index of Multiple Deprivations 2007, which is an aggregate score for each English catchment area. Subjects were linked to their corresponding catchment area by postcode of residence and associations with deprivation were analyzed in quintiles, with quintile 1 being the most deprived.

Colorectal cancer variables

CRC site was classified based on the first CRC coding episode into right sided, left sided, and unspecified (*appendix 3*). Coding records of initially unspecified site CRC were examined and if a more specific code had been used subsequently, this was used to determine the CRC site. Colonic polyps were identified from ICD-10 codes (*appendix 4*).

Distant metastases were identified by ICD-10 codes (*appendix 5*) up to 12 months from diagnosis date and were used as a surrogate marker of CRC stage at diagnosis, as Dukes' staging is not recorded in HES. Codes for metastases can occasionally be miscoded as a primary neoplasm (eg, lung), and therefore primary malignancy codes were also used, provided that they were recorded in the 12 months subsequent to CRC diagnosis (*appendix 5*). Surgery and chemotherapy were identified by respective OPCS-4 codes (*appendix 6*).

Survival analysis

Survival analysis adjusted for gender, age, deprivation, and co-morbidity was calculated from the CRC diagnosis date of PCCRC cases and controls using date of death from ONS. Subjects who were not diagnosed by colonoscopy were not included to avoid potential lead time bias due to the method of determining date of diagnosis from HES.

Provider variables

For the purpose of this study, all endoscopy units operating within the same NHS organisation were analysed as a single provider. Individual providers were stratified by colonoscopy volume, bowel cancer screening program (BCSP) status and the percentage of CRCs diagnosed during an emergency rather than an elective episode to determine if there was an association with PCCRC. Colonoscopy volume was determined from the total number of colonoscopies performed during the study period at each provider and separated into tertiles. A BCSP accredited provider had at least one endoscopy unit accredited with BCSP status by the end of the study period. The percentage of CRCs diagnosed as an emergency at a provider was the ratio of CRCs diagnosed during an acute (unplanned) admission divided by all CRCs, including CRCs diagnosed during an elective episode.

Ethics

As only pseudonymized information was used in this study, ethics approval was not necessary. HES data are available under a data-sharing agreement for the purposes of service evaluation.

Statistical methodology

Statistical analysis was carried out with STATA SE v13.1 (Statacorp LP, Tex, USA). Analysis of variance and χ^2 tests were used to compare differences in continuous and categorical variables respectively. Associations with PCCRC were examined by univariate and multivariate logistic regression. A multivariate model was constructed to determine associations with PCCRC after adjusting for gender, age, Charlson co-morbidity index, procedure type (emergency or elective), CRC site (left side of colon or right side of colon), metastases, and procedure year. For tests of significance, p values <0.05 were considered significant. All odds ratios, 95% confidence intervals, and associated p values are the result of multivariate analysis unless stated otherwise. Unadjusted Kaplan-Meier analysis and Cox proportional hazards modeling after adjustment for gender, age, deprivation, and co-morbidity were used to compare survival in PCCRC cases and controls.

Results

Study cohort

Between April 2003 and March 2009, 1,439,684 colonoscopies were identified and 67,202 subjects were diagnosed with CRC during this period. Out of the 67,202 CRC subjects, there were 8147 (12.1%) PCCRC subjects: 1796 (2.7%) PCCRC 6 to 12 months; 3772 (5.6%) PCCRC 12 to 36 months, and 2579 (3.8%) PCCRC 36 to 60 months. A total of 59,055 CRC subjects had not had a colonoscopy between 6 and 60 months before CRC diagnosis and served as controls. Overall, 0.66% or 1 in every 150 subjects developed PCCRC after a colonoscopy that did not diagnose CRC.

Validation of colonoscopy and colorectal cancer populations

The total number of colonoscopies carried out between 2007 and 2010 at UHB was 8708 and 8292 colonoscopies (95.2%) were coded in HES for UHB for the equivalent 4-year period. The CRC population was validated by comparing CRC cases recorded in HES (315,515) to CRC cases reported from 2002 to 2011 by NCIN (312,984)[14], showing a concordance of over 99%. The CRC population was further validated by comparing the 70.4% surgical rate for CRC from HES with the National Bowel Cancer Audit, which reported that 75.7% of CRC patients enrolled in the audit underwent surgery between 2008 and

2011.[15, 16, 17] All of the validation processes showed a good correlation between HES data and independent data sources, suggesting the study methodology was valid.

Subject characteristics

The characteristics of cases with PCCRC and CRC controls are shown in Table 1. PCCRC subjects (mean age 70.7 ± 11.4 years) were older than controls (mean age 70.2 ± 11.4 years)($p < 0.001$). The risk of PCCRC appeared to increase with age on univariate analysis, but only subjects aged 70 to 74 were associated with PCCRC compared with subjects under 60, after adjusting for confounding factors. PCCRC subjects were more likely to be female. Subjects with the most co-morbidities (Charlson co-morbidity index of 5 or greater) were associated with PCCRC. PCCRC was not associated with differences in ethnicity or deprivation.

Colonoscopy variables and findings

The influence of colonoscopy variables and findings on PCCRC are shown in Table 2. The majority of CRC were diagnosed during an elective colonoscopy. However, being diagnosed during an emergency colonoscopy reduced the risk of PCCRC nearly by half. There was minor increased risk of PCCRC on univariate analysis in colonoscopies carried out at the weekend compared with during the week.

PCCRC was more likely to be associated with CRC in the right side of the colon. Colonic polyps were coded in 21.6% of the colonoscopies that did not detect CRC in the PCCRC group. Polypectomy was coded in a further 18.9%. On univariate analysis, this was higher than both the recorded polyp rate of 9.8% (2.52 (95% CI, 2.39-2.65), $p < 0.0001$) and polypectomy rate of 11.3% (1.82 (95% CI, 1.72-1.92), $p < 0.0001$) from all colonoscopies during the study period. Furthermore, the polyp and polypectomy rates were both higher in the PCCRC 6 to 12 months group on univariate analysis, than in the PCCRC 12 to 36 months ($p < 0.0001$), and PCCRC 36 to 60 months ($p < 0.0001$) groups.

Colorectal outcomes and survival

The prevalence of metastatic disease within 12 months of CRC diagnosis in PCCRC cases and controls are shown in Table 3. PCCRC cases were up to twice as likely to be diagnosed with

lung, peritoneal, and bone metastases within 12 months of CRC diagnosis. However, lymph node metastases were more prevalent in controls than PCCRC cases, suggesting coding bias related to the increased rate of surgery in control subjects described later.

On univariate analysis, PCCRC cases were less likely to undergo surgery compared with controls (0.33 (95% CI, 0.32-0.35), $p<0.0001$) or chemotherapy (0.66 (95% CI, 0.62-0.69), $p<0.0001$). Overall survival was also worse in PCCRC subjects compared with controls, with a median survival of 5.8 years in controls compared with 2.1 years in the PCCRC 6- to 12-month group, 2.0 years in the PCCRC 12- to 36-month group, and 3.5 years in the PCCRC 36- to 60-month group (figure 1). after adjusting for age, gender, co-morbidity, and deprivation, survival outcomes remained worse for PCCRC subjects with a hazard ratio of 1.17 (95% CI, 1.10- 1.24)($p<0.0001$), 1.26 (95% CI, 1.20-1.31)($p<0.0001$) and 1.20 (95% CI, 1.13- 1.27)($p<0.0001$) for the PCCRC 6 to 12 months, PCCRC 12 to 36 months, and PCCRC 36 to 60 months respectively when compared with controls.

Individual provider variables

The influence of provider variables on PCCRC are shown in Table 4. There was a more than twofold variation in PCCRC rates between individual providers in England during the study period (figure 2). On univariate analysis, medium colonoscopy volume providers and low volume providers were both more likely to be associated with PCCRC than high volume providers. after adjusting for other variables in the multivariate model an association with medium volume providers remained. BCSP accreditation status and the percentage of CRC diagnosed as an emergency were not associated with an increased risk of PCCRC.

PCCRC rates over time

The number of colonoscopies recorded in HES has increased by almost 2-fold over the study period. Despite the increase in colonoscopy numbers performed year on year, the annual rate of PCCRC has steadily fallen over the study period ($p<0.0001$)(figure 3). The annual PCCRC rate decreased from 13.8% in 2003 to 2004 to 11.9% by the end of study period in 2008 to 2009 with the reduction seen mainly in the PCCRC 6- to 12-month and PCCRC 12- to 36-month groups.

Discussion

The overall PCCRC rate of 12.1% in 67,202 subjects in England between 2003 and 2009 appears higher than previously published figures. However, some previous studies have calculated the PCCRC rate by only including CRC subjects with a colonoscopy up to 36 months before diagnosis and the comparable figure from the present study is 8.3%. A Canadian study of 14,064 CRC subjects reported a PCCRC rate of 9.0% between 2000 and 2005.[12] Using the Surveillance, Epidemiology, and End Results (SEER)-Medicare database in the USA, a PCCRC rate of 7.2% was reported between 1994 to 2005 from a study of 57,839 CRC subjects.[19] A further population based study from Utah, USA, with 2659 CRC subjects between 1995 and 2009 described a PCCRC rate of 6% when subjects with a colonoscopy up to 60 months before CRC diagnosis were included.[4] In Europe, 2 recent studies have reported much lower PCCRC rates. A Danish population based study between 2000 to 2009 included 37,044 CRC subjects and concluded that only 2.7% of CRC subjects have had a colonoscopy that failed to diagnose CRC 1 to 5 years before diagnosis.[5] A second study from the Netherlands analyzed 5107 CRC subjects between 2001 to 2010 from three providers and found a PCCRC rate of only 2.9% for subjects with a colonoscopy up to 60 months before CRC diagnosis.[20] In addition to potential variations in subject and colonoscopy factors between the difference studies, the wide range of reported PCCRC rates are likely to be contributed to by methodological differences.[13]

In the present study, PCCRC was associated with older subjects, female gender, an increased number of co-morbidities and right-sided CRC, which is in keeping with findings from other studies of PCCRC. [3, 12, 19, 21] The association between increasing age and PCCRC was less marked on multivariate analysis and this may relate to confounding from increasing co-morbidity in the elderly. Elderly patients are more likely to have inadequate bowel preparation, thus reducing mucosal visualisation and detection of polyps and early CRC.[22, 23] Female patients are more likely to have had previous abdominal and pelvic surgery, which may increase the technical difficulty of colonoscopy and impair patient tolerance, reducing the cecal intubation rate.[24] In addition to factors that have an adverse effect on cecal intubation rate, right-sided CRC are more likely to arise from flat, non-polypoid

adenomatous lesions[20, 25] that poor bowel preparation may make difficult to detect. This will contribute to the association of right-sided CRC with PCCRC.

Over a fifth of PCCRC subjects had colonic polyps or polypectomy coded during the most recent colonoscopy before CRC diagnosis. This is higher than the average polypectomy rate in all colonoscopy procedures during the same period. Furthermore, polyp and polypectomy coding rates were highest in the PCCRC 12- to 36-month group. Prior polypectomy has been reported to double the risk of PCCRC[19], with up to 19% of CRCs occurring in the same anatomic segment as a previously resected adenoma.[8] Paradoxically, colonoscopists with higher polypectomy rates have been reported to be associated with a lower risk of PCCRC[12, 19], presumably as they detect more polyps and remove them more completely than other colonoscopists. Incomplete polypectomy, or inadequate biopsy sampling of polyps, is therefore a key modifiable risk factor for PCCRC and ensuring adequate follow-up and assessment after polypectomy may reduce PCCRC rates.

PCCRC subjects appeared to have worse outcomes in terms of both treatment after diagnosis (surgery and chemotherapy) and overall survival. Previous studies have reported no survival difference between PCCRC subjects and controls[5, 21] with one recent study even reporting a survival benefit in the PCCRC subjects, which was likely to be due to earlier CRC stage at diagnosis in the PCCRC subjects.[4] In the current study, PCCRC subjects were older, had greater co-morbidities and were more likely to present with distant metastases within 12 months of diagnosis compared with controls. All these factors contributed to the reduced rates of curative surgery or palliative chemotherapy for PCCRC subjects and will have contributed to worse survival. Adjusting the survival analyses for differences in ages, gender, co-morbidity and deprivation still revealed worse survival for PCCRC subjects and, at least in England, PCCRC is clearly associated with worse survival. Survival in PCCRC subjects would have been potentially better if earlier opportunities to diagnose their CRC had been taken.

Previous studies have reported that PCCRC was not associated with endoscopist procedure volume[12] and that higher colonoscopy volumes may even be positively associated with PCCRC surprisingly.[19] In the current study, there was a large variation in PCCRC rates

1 between individual providers across England but PCCRC appeared to be associated with
2 lower colonoscopy volume providers. This result should be interpreted with caution. We did
3 not have access to colonoscopy quality indicators such as cecal intubation and adenoma
4 detection rates that are likely to be potentially more important factors in PCCRC incidence.
5
6
7

8
9 Colonoscopy undertaken during an emergency admission covered 10% of procedures
10 examined and was associated with a lower risk of PCCRC at 9% compared with 14% for
11 elective procedures. Patients presenting as an emergency may have more advanced
12 colorectal cancer and therefore a lower chance of PCCRC.
13
14
15
16
17

18
19 The annual PCCRC rate in England has fallen steadily over the study period from 13.8% to
20 11.9%, at least partly due to improving colonoscopy standards over the corresponding time
21 period. In 2003, a multi-regional audit in England including 9223 colonoscopies reported
22 that mean cecal intubation rate was only 76.9%.[26] A subsequent national audit in 2011 of
23 20085 colonoscopies found that the cecal intubation rate had improved to 92.3%.[1] The
24 PCCRC rate is likely to continue to improve in recent years given changes in colonoscopy
25 practice, including the recognition of the importance of minimum withdrawal times [27],
26 bowel preparation improvements[28], and better endoscopic recognition of sessile serrated
27 polyps[25], subsequent to the study period.
28
29
30
31
32
33
34
35
36
37

38
39 The use of a national hospital dataset enabled us to undertake one of the largest PCCRC
40 studies to date, including the vast majority of colonoscopies performed during a period of
41 rising colonoscopy standards. The quality of diagnostic and procedural coding in HES has
42 been previously investigated and there was a high concordance when compared with
43 independent national data sources.[1, 10, 29] However, we did not have the opportunity to
44 link our HES dataset directly to cancer registry data due to restrictions under which the data
45 is held and therefore, in order to validate the methodology chosen, colonoscopy and CRC
46 populations were compared with national cancer databases and a local data sample and
47 revealed a good correlation. The completeness and accuracy of coding in HES is still a
48 potential source of concern. For example, the diagnosis date may not be recorded
49 accurately in HES due to the need for histological confirmation before CRC coding and
50 therefore a colonoscopy within 6 months of CRC coding had to be considered the diagnostic
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

1 procedure. There are also limitations in HES concerning coding of colonoscopy procedures,
2 polyps, polypectomy, presence of metastases, surgery and chemotherapy and the figures
3 included may be an over or under estimate, though this is likely to affect PCCRC cases and
4 controls equally. A further limitation is that key procedure information such as the bowel
5 preparation quality, sedation doses, colonoscopist grade and specialty, extent of
6 examination, completeness of polypectomy, and number of biopsy specimens taken are not
7 recorded in HES and all may influence the PCCRC risk. Furthermore, due to the HES coding
8 hierarchy, indication, presence of diverticular disease and history of abdominal or pelvic
9 surgery may not be coded, partly due to under-reporting by colonoscopists when significant
10 pathology or CRC are found and again each may be important risk factors for PCCRC. As HES
11 does not record polyp histology or the International Classification of Diseases for Oncology
12 (ICD-O) codes, the lack of data on polyp and CRC histology and Duke's staging further limits
13 analysis of potential causes of PCCRC (de novo CRC, incomplete adenoma resection, missed
14 lesion or biopsy failed to detect CRC), and survival in PCCRC subjects.
15
16
17
18
19
20
21
22
23
24
25
26
27
28

29 In conclusion, the PCCRC rate was 12.1% in England between 2003 and 2009. PCCRC was
30 associated with older age, female gender, increasing co-morbidity, procedure related
31 factors (elective procedures and right-sided CRC), and provider colonoscopy volume.
32
33
34

35 Despite the encouraging fall in annual PCCRC rate over the study period, the PCCRC rate
36 should be a routinely measured endoscopy unit colonoscopy quality marker, and potentially
37 avoidable risk factors for PCCRC should be addressed.
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

Table 1. The characteristics of post-colonoscopy colorectal cancer cases and controls

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	P value	Odds ratio	95% CI	P value
Total subjects (number)						Univariate			Multivariate		
	1796 (2.7)	3772 (5.6)	2579 (3.8)	8147 (12.1)	59055 (87.9)	-	-	-	-	-	-
Mean age \pmSD (years)											
	71.5 \pm 11.4	70.9 \pm 11.7	69.8 \pm 10.8	70.7 \pm 11.4	70.2 \pm 11.4	-	-	<0.001			
Age group (number (%))											
Under 60	263 (3.2)	598 (7.3)	415 (5.1)	1276 (15.7)	9849 (16.7)	Ref					
60-64	167 (2.0)	367 (4.5)	288 (3.5)	822 (10.1)	6749 (11.4)	0.94	0.86-1.03	0.1928	0.95	0.86-1.04	0.277
65-69	217 (2.7)	531 (6.5)	435 (5.3)	1183 (14.5)	8810 (14.9)	1.04	0.95-1.13	0.4044	1.03	0.94-1.12	0.537
70-74	344 (4.2)	648 (8.0)	488 (6.0)	1480 (18.2)	10229 (17.3)	1.12	1.03-1.21	0.0067	1.09	1.00-1.18	0.039
75-79	359 (4.4)	678 (8.3)	499 (6.1)	1536 (18.9)	10698 (18.1)	1.11	1.02-1.20	0.0109	1.07	0.98-1.16	0.159
80+	446 (5.5)	950 (11.7)	454 (5.6)	1850 (22.7)	12720 (21.5)	1.12	1.04-1.21	0.0029	1.08	1.00-1.17	0.065
Gender (number (%))											
Male	974 (12.0)	1974 (24.2)	1340 (16.4)	4288 (52.6)	33057 (56.0)	Ref	-	-	Ref	-	-
Female	822 (10.1)	1798 (22.1)	1239 (15.2)	3859 (47.4)	25998 (44.0)	1.14	1.09-1.20	<0.0001	1.13	1.08-1.19	<0.001
Charlson co-morbidity index (number (%))											
0	1514 (18.6)	3210 (39.4)	2235 (27.4)	6959 (85.4)	50663 (85.8)	Ref	-	-	Ref	-	-
1-4	154 (1.9)	298 (3.7)	210 (2.6)	662 (8.1)	4764 (8.1)	1.01	0.93-1.10	0.7896	1.06	0.97-1.16	0.195
5+	128 (1.6)	264 (3.2)	134 (1.6)	526 (6.5)	3628 (6.1)	1.06	0.96-1.16	0.2641	1.16	1.05-1.28	0.003
Deprivation quintile (number (%))											
1 (most)	329 (4.0)	637 (7.8)	393 (4.8)	1359 (16.7)	10015 (17.0)	Ref	-	-	-	-	-
2	365 (4.5)	740 (9.1)	499 (6.1)	1604 (19.7)	11258 (19.1)	1.05	0.97-1.13	0.2153	-	-	-
3	333 (4.1)	782 (9.6)	551 (6.8)	1666 (20.4)	12399 (21.0)	0.99	0.91-1.07	0.8002	-	-	-
4	387 (4.8)	784 (9.6)	568 (7.0)	1739 (21.3)	12642 (21.4)	1.01	0.94-1.09	0.7242	-	-	-
5 (least)	381 (4.7)	823 (10.1)	566 (6.9)	1770 (21.7)	12620 (21.4)	1.03	0.96-1.11	0.3905	-	-	-
Ethnicity (number (%))											
White	1656 (20.3)	3536 (43.4)	2467 (30.3)	7659 (94.0)	54512 (92.3)	Ref	-	-	-	-	-
Asian	21 (0.3)	55 (0.7)	36 (0.4)	112 (1.4)	788 (1.3)	1.01	0.83-1.23	0.9097	-	-	-

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49

Afro-Caribbean	25 (0.3)	53 (0.7)	27 (0.3)	105 (1.3)	823 (1.4)	0.91	0.74-1.11	0.3553	-	-	-
Chinese	0	0	0	12 (0.1)	118 (0.2)	0.72	0.40-1.30	0.2865	-	-	-
Mixed	0	0	0	18 (0.2)	160 (0.3)	0.80	0.49-1.30	0.3719	-	-	-
Others	12 (0.1)	21 (0.3)	21 (0.3)	54 (0.7)	341 (0.6)	1.13	0.85-1.50	0.4156	-	-	-
Unknown	74 (0.9)	95 (1.2)	18 (0.2)	187 (2.3)	2313 (3.9)	0.58	0.49-0.67	<0.0001	-	-	-

Odds ratios with 95% confidence intervals and p values for PCCRC (all) compared with controls
PCCRC – post-colonoscopy colorectal cancer

Table 2. The colonoscopy characteristics and findings of post-colonoscopy colorectal cancer cases and controls

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	P value	Odds ratio	95% CI	P value
Procedure day (number (%))						Univariate			Multivariate		
Weekday	1736 (21.3)	3628 (44.5)	2486 (30.5)	7850 (96.4)	57249 (96.9)	Ref	-	-	-	-	-
Weekend	60 (0.7)	144 (1.8)	93 (1.1)	297 (3.6)	1806 (3.1)	1.19	1.06-1.36	0.0044	-	-	-
Procedure type (number (%))											
Elective	1622 (19.9)	3473 (42.6)	2455 (30.1)	7550 (92.7)	52605 (89.1)	Ref	-	-	Ref	-	-
Emergency	174 (2.1)	299 (3.7)	124 (1.5)	597 (7.3)	6450 (10.9)	0.64	0.59-0.70	<0.0001	0.54	0.59-0.69	<0.0001
Colorectal cancer location (number (%))											
Left sided	897 (11.0)	1754 (21.5)	1260 (15.5)	3911 (48.0)	34703 (58.8)	Ref	-	-	Ref	-	-
Right sided	535 (6.6)	1242 (15.2)	919 (11.3)	2696 (33.1)	20751 (35.1)	1.15	1.09-1.21	<0.0001	1.17	1.11-1.23	<0.0001
Unknown/overlapping sites	364 (4.5)	776 (9.5)	400 (4.9)	1540 (18.9)	3601 (6.1)	3.79	3.54-4.06	<0.0001	3.72	3.46-3.99	<0.0001
Polyp/ polypectomy coded (number (%))											
Polyp coded	491 (6.0)	742 (9.1)	523 (6.4)	1756 (21.6)	141799* (9.8)	2.52⁺	2.39-2.65⁺	<0.0001⁺	-	-	-
No polyp coded	1305 (16.0)	3030 (37.2)	2056 (25.2)	6391 (78.4)	1300714* (90.2)	Ref	-	-	-	-	-
Polypectomy coded	348 (4.3)	669 (8.2)	523 (6.4)	1540 (18.9)	162364* (11.3)	1.82⁺	1.72-1.92⁺	<0.0001⁺	-	-	-
No polypectomy coded	1448 (17.8)	3103 (38.1)	2056 (25.2)	6607 (81.1)	1280150* (89.7)	Ref	-	-	-	-	-

Odds ratios with 95% confidence intervals and p values for all PCCRC compared with controls

PCCRC – post-colonoscopy colorectal cancer

* From all colonoscopies

+ Univariate analysis comparing all PCCRC with all colonoscopies during study period.

Table 3. The prevalence of metastases within 12 months of colorectal cancer diagnosis in post-colonoscopy colorectal cancer cases and controls

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	p value	Odds ratio	95% CI	P value
Subjects with metastases within 12 months of diagnosis (number (%))						Univariate			Multivariate		
Liver metastases	276 (3.4)	619 (7.6)	365 (4.5)	1260 (15.5)	8545 (14.5)	1.08	1.01-1.15	0.017	0.97	0.91-1.05	0.486
Lung metastases	154 (1.9)	345 (4.2)	182 (2.2)	681 (8.4)	3104 (5.3)	1.64	1.51-1.79	<0.0001	1.61	1.46-1.77	<0.0001
Peritoneal metastases	75 (0.9)	166 (2.0)	102 (1.3)	343 (4.2)	1903 (3.2)	1.32	1.17-1.48	<0.0001	1.27	1.12-1.44	<0.0001
Bone metastases	45 (0.6)	106 (1.3)	78 (1.0)	229 (2.8)	678 (1.1)	2.49	2.14-2.90	<0.0001	2.21	1.88-2.60	<0.0001
Lymph node metastases	136 (1.7)	282 (3.5)	231 (2.8)	649 (8.0)	6459 (10.9)	0.70	0.65-0.76	<0.0001	0.75	0.69-0.82	<0.0001
Treatment outcome after diagnosis (number (%))											
Surgery	791 (9.7)	1661 (20.4)	1337 (16.4)	3789 (46.5)	42790 (72.5)	0.33	0.32-0.35	<0.0001	-	-	-
Chemotherapy	422 (5.2)	911 (11.2)	594 (7.3)	1927 (23.7)	18908 (32.0)	0.66	0.62-0.69	<0.0001	-	-	-

Odds ratios with 95% confidence intervals and *P* values for PCCRC (all) compared with controls.

PCCRC – post-colonoscopy colorectal cancer.

Table 4. The influence of provider variables on post-colonoscopy colorectal cancer

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	p value	Odds ratio	95% CI	P value
Colonoscopy volume by NHS provider (number (%))						Univariate			Multivariate		
High volume providers (>1680 pa)	955 (11.7)	1993 (24.5)	1415 (17.4)	4363 (53.6)	33353 (56.5)	Ref	-	-	Ref	-	-
Medium volume providers	704 (8.6)	1486 (18.2)	994 (12.2)	3184 (39.1)	21942 (37.2)	1.11	1.06-1.16	<0.0001	1.13	1.01-1.27	0.035
Low-volume providers (<747 pa)	137 (1.7)	293 (3.6)	170 (2.1)	600 (7.4)	3760 (6.4)	1.22	1.11-1.34	<0.0001	1.05	0.98-1.12	0.161
BCSP status (number (%))											
BCSP provider	959 (11.8)	2064 (25.3)	1396 (17.1)	4419 (54.2)	31780 (53.8)	Ref	-	-	-	-	-
Non-BCSP provider	837 (10.3)	1708 (21.0)	1183 (14.5)	3728 (45.8)	27275 (46.2)	0.98	0.94-1.03	0.4690	0.96	0.90-1.03	0.255
Percentage of CRC diagnosed during an emergency admission by NHS provider (number (%))											
Low-percentage providers (<27.3%)	408 (5.0)	848 (10.4)	629 (7.7)	1885 (23.1)	14270 (24.2)	0.91	0.84-0.98	0.0115	0.96	0.87- 1.06	0.443
Medium percentage providers	1068 (13.1)	2273 (27.9)	1530 (18.8)	4871 (59.8)	35211 (59.6)	0.95	0.89-1.01	0.1299	0.96	0.85-1.09	0.531
High-percentage providers (>33.9%)	320 (3.9)	651 (8.0)	420 (5.2)	1391 (17.1)	9572 (16.2)	Ref	-	-	Ref	-	-

Odds ratios with 95% confidence intervals and p values for PCCRC (all) compared with controls

PCCRC – post-colonoscopy colorectal cancer

BCSP – Bowel Cancer Screening Program

Figure 1. Post-colonoscopy colorectal cancer rates by individual provider in England between 2003 and 2009.

Figure 2. Unadjusted survival after colorectal cancer diagnosis in post-colonoscopy colorectal cancer cases and control subjects.

Figure 3 Post-colonoscopy colorectal cancer rates and colonoscopy volume in England by year.

Appendix 1 - OPCS-4 codes for colonoscopy

H20.1 Snare polypectomy

H20.6 Polypectomy with colonoscopy

H22.1 Diagnostic fibreoptic endoscopic examination of colon and biopsy of lesion of colon

H22.8 Other specified diagnostic endoscopic examination of colon

H22.9 Unspecified diagnostic endoscopic examination of colon

Appendix 2 - ICD-10 codes for colorectal cancers

C18 Malignant neoplasm of colon - excluding C18.1 (malignant neoplasm of appendix)

C19 Malignant neoplasm of rectosigmoid junction

C20 Malignant neoplasm of rectum

Appendix 3 – ICD-10 codes for colorectal cancer (CRC) sites

Right sided CRC

C18.0 Caecum, Ileocaecal valve

C18.2 Ascending colon

C18.3 Hepatic flexure

C18.4 Transverse colon

Left sided CRC

C18.5 Splenic flexure

C18.6 Descending colon

C18.7 Sigmoid colon

C19 Rectosigmoid junction

C20 Rectum

Unspecified CRC location

C18.8 Overlapping lesion of colon

C18.9 Colon, unspecified

Appendix 4 - ICD-10 codes for colorectal polyps

D12.0 Caecal polyp(s)

1 D12.2 Ascending colon polyp(s)

2 D12.3 Transverse colon, hepatic flexure, splenic flexure polyp(s)

3 D12.4 Descending colon polyp(s)

4 D12.5 Sigmoid colon polyp(s)

5 D12.6 Colon, site unspecified polyp(s)

6 D12.7 Rectosigmoid junction polyp(s)

7 D12.8 Rectal polyp(s)

8 **Appendix 5 - ICD-10 codes for metastases**

9 C77.1 Intrathoracic lymph nodes

10 C77.2 Intra-abdominal lymph nodes

11 C77.4 Inguinal and lower limb lymph nodes

12 C77.5 Intrapelvic lymph nodes

13 C78.0 Secondary malignant neoplasm of lung

14 C78.6 Secondary malignant neoplasm of retroperitoneum and peritoneum

15 C78.7 Secondary malignant neoplasm of liver and intrahepatic bile duct

16 C79.5 Secondary malignant neoplasm of bone and bone marrow

17 C34 Malignant neoplasm of bronchus and lung

18 C48 Malignant neoplasm of retroperitoneum and peritoneum

19 C22 Malignant neoplasm of liver

20 C40-C41 Malignant neoplasms of bone and articular cartilage

21 **Appendix 6- OPCS-4 codes for surgical procedures**

22 H04 Total excision of colon and rectum

23 H05 Total excision of colon

24 H06 Extended excision of right hemicolon

25 H07 Other excision of right hemicolon

26 H08 Excision of transverse colon

27 H09 Excision of left hemicolon

28 H10 Excision of sigmoid colon

29 H11 Other excision of colon

30 H29 Subtotal excision of colon

1 H33 Excision of rectum
2 H40 Operations on rectum through anal sphincter
3
4 H122 Excision of lesion of colon NEC
5
6 H123 Destruction of lesion of colon NEC
7
8 H128 Other specified extirpation of lesion of colon
9
10 H129 Unspecified extirpation of lesion of colon
11
12 H341 Open excision of lesion of rectum
13
14 H345 Open destruction of lesion of rectum
15
16 H348 Other specified open extirpation of lesion of rectum
17
18 H349 Unspecified open extirpation of lesion of rectum
19
20 H402 Trans-sphincteric excision of lesion of rectum
21
22 H403 Trans-sphincteric destruction of lesion of rectum
23
24 OPCS-4 codes for chemotherapy
25
26 X70 Procurement of drugs for chemotherapy for neoplasm in Bands 1-5
27
28 X71 Procurement of drugs for chemotherapy for neoplasm in Bands 6-10
29
30 X72 Delivery of Chemotherapy for neoplasm
31
32 X73 Delivery of oral chemotherapy for neoplasm
33
34 X352 Intravenous chemotherapy
35
36 X384 Subcutaneous chemotherapy
37
38 X373 Intramuscular chemotherapy
39
40 Z082 Follow up examination after chemotherapy for malignant neoplasm
41
42 Z511 Chemotherapy session for neoplasm
43
44 Z542 Convalescence following chemotherapy
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

References

- 1 Gavin DR, Valori RM, Anderson JT, Donnelly MT, Williams JG, Swarbrick ET. The national colonoscopy audit: a nationwide assessment of the quality and safety of colonoscopy in the UK. *Gut* 2013;**62**:242-9.
- 2 Coleman MP, Forman D, Bryant H, Butler J, Rachet B, Maringe C, *et al.* Cancer survival in Australia, Canada, Denmark, Norway, Sweden, and the UK, 1995-2007 (the International Cancer Benchmarking Partnership): an analysis of population-based cancer registry data. *Lancet* 2011;**377**:127-38.
- 3 Bressler B, Paszat LF, Chen Z, Rothwell DM, Vinden C, Rabeneck L. Rates of new or missed colorectal cancers after colonoscopy and their risk factors: a population-based analysis. *Gastroenterology* 2007;**132**:96-102.
- 4 Samadder NJ, Curtin K, Tuohy TM, Pappas L, Boucher K, Provenzale D, *et al.* Characteristics of missed or interval colorectal cancer and patient survival: a population-based study. *Gastroenterology* 2014;**146**:950-60.
- 5 Erichsen R, Baron JA, Stoffel EM, Laurberg S, Sandler RS, Sorensen HT. Characteristics and survival of interval and sporadic colorectal cancer patients: a nationwide population-based cohort study. *The American journal of gastroenterology* 2013;**108**:1332-40.
- 6 Arain MA, Sawhney M, Sheikh S, Anway R, Thyagarajan B, Bond JH, *et al.* CIMP status of interval colon cancers: another piece to the puzzle. *The American journal of gastroenterology* 2010;**105**:1189-95.
- 7 Sawhney MS, Farrar WD, Gudiseva S, Nelson DB, Lederle FA, Rector TS, *et al.* Microsatellite instability in interval colon cancers. *Gastroenterology* 2006;**131**:1700-5.
- 8 Robertson DJ, Lieberman DA, Winawer SJ, Ahnen DJ, Baron JA, Schatzkin A, *et al.* Colorectal cancers soon after colonoscopy: a pooled multicohort analysis. *Gut* 2014;**63**:949-56.
- 9 Health & Social Care Information Centre. *Hospital Episode Statistics*; www.hscic.gov.uk/hes [Accessed 29 December 2013].
- 10 Shaihi M, Thompson E, Kapoor N, Powell G, Sturges RP, Stern N, *et al.* Variation in gastroscopy rate in English general practice and outcome for oesophagogastric cancer: retrospective analysis of Hospital Episode Statistics. *Gut* 2014;**63**:250-61.
- 11 Weller D, Vedsted P, Rubin G, Walter FM, Emery J, Scott S, *et al.* The Aarhus statement: improving design and reporting of studies on early cancer diagnosis. *British journal of cancer* 2012;**106**:1262-7.
- 12 Baxter NN, Sutradhar R, Forbes SS, Paszat LF, Saskin R, Rabeneck L. Analysis of administrative data finds endoscopist quality measures associated with postcolonoscopy colorectal cancer. *Gastroenterology* 2011;**140**:65-72.
- 13 Morris EJ, Rutter MD, Finan PJ, Thomas JD, Valori R. Post-colonoscopy colorectal cancer (PCCRC) rates vary considerably depending on the method used to calculate them: a retrospective observational population-based study of PCCRC in the English National Health Service. *Gut* 2015;**64**:1248-56.
- 14 National Cancer Intelligence Network. UK Cancer e-Atlas by cancer networks.
- 15 Health & Social Care Information Centre. *National Bowel Cancer Audit 2009*; www.hscic.gov.uk/catalogue/PUB02587/nati-clin-audi-supprog-bowe-canc-2009-rep2.pdf [Accessed 2 June 2014]. 2009.
- 16 Health & Social Care Information Centre. *National Bowel Cancer Audit 2010*; www.hscic.gov.uk/catalogue/PUB02586/nati-clin-audi-supprog-bowe-canc-2010-rep1.pdf [Accessed 2 June 2014]. 2010.
- 17 Health & Social Care Information Centre. *National Bowel Cancer Audit 2011*; www.hscic.gov.uk/catalogue/PUB02576/nati-clin-audi-supprog-bowe-canc-2011-rep1.pdf [Accessed 2 June 2014]. 2011.

- 18 Charlson ME, Pompei P, Ales KL, MacKenzie CR. A new method of classifying prognostic comorbidity in longitudinal studies: development and validation. *Journal of chronic diseases* 1987;**40**:373-83.
- 19 Cooper GS, Xu F, Barnholtz Sloan JS, Schluchter MD, Koroukian SM. Prevalence and predictors of interval colorectal cancers in medicare beneficiaries. *Cancer* 2012;**118**:3044-52.
- 20 le Clercq CM, Bouwens MW, Rondagh EJ, Bakker CM, Keulen ET, de Ridder RJ, *et al*. Postcolonoscopy colorectal cancers are preventable: a population-based study. *Gut* 2014;**63**:957-63.
- 21 Singh H, Nugent Z, Demers AA, Bernstein CN. Rate and predictors of early/missed colorectal cancers after colonoscopy in Manitoba: a population-based study. *The American journal of gastroenterology* 2010;**105**:2588-96.
- 22 Romero RV, Mahadeva S. Factors influencing quality of bowel preparation for colonoscopy. *World journal of gastrointestinal endoscopy* 2013;**5**:39-46.
- 23 Hong SN, Sung IK, Kim JH, Choe WH, Kim BK, Ko SY, *et al*. The Effect of the Bowel Preparation Status on the Risk of Missing Polyp and Adenoma during Screening Colonoscopy: A Tandem Colonoscopic Study. *Clinical endoscopy* 2012;**45**:404-11.
- 24 Shah HA, Paszat LF, Saskin R, Stukel TA, Rabeneck L. Factors associated with incomplete colonoscopy: a population-based study. *Gastroenterology* 2007;**132**:2297-303.
- 25 Tadros M, Anderson JC. Serrated polyps: clinical implications and future directions. *Current gastroenterology reports* 2013;**15**:342.
- 26 Bowles CJ, Leicester R, Romaya C, Swarbrick E, Williams CB, Epstein O. A prospective study of colonoscopy practice in the UK today: are we adequately prepared for national colorectal cancer screening tomorrow? *Gut* 2004;**53**:277-83.
- 27 Simmons DT, Harewood GC, Baron TH, Petersen BT, Wang KK, Boyd-Enders F, *et al*. Impact of endoscopist withdrawal speed on polyp yield: implications for optimal colonoscopy withdrawal time. *Alimentary pharmacology & therapeutics* 2006;**24**:965-71.
- 28 Aoun E, Abdul-Baki H, Azar C, Mourad F, Barada K, Berro Z, *et al*. A randomized single-blind trial of split-dose PEG-electrolyte solution without dietary restriction compared with whole dose PEG-electrolyte solution with dietary restriction for colonoscopy preparation. *Gastrointestinal endoscopy* 2005;**62**:213-8.
- 29 Moller H, Richards S, Hanchett N, Riaz SP, Luchtenborg M, Holmberg L, *et al*. Completeness of case ascertainment and survival time error in English cancer registries: impact on 1-year survival estimates. *British journal of cancer* 2011;**105**:170-6.

***Acronyms (list all acronyms used in paper with their spell-outs)**

Bowel cancer screening program (BCSP)

Colorectal cancers (CRC)

Hospital Episode Statistics (HES)

International Classification of Diseases version 10 (ICD-10)

National Cancer Intelligence Network (NCIN)

National Health Service (NHS)

Office for National Statistics (ONS)

Office of Population Censuses and Surveys Classification of Interventions and Procedures 4th revision (OPCS-4)

Surveillance, Epidemiology, and End Results Medicare database (SEER)

Post-colonoscopy colorectal cancer (PCCRC)

University Hospital Birmingham (UHB)



Journal CME Conflict of Interest: Disclosure and Attestation

Lead Author: Dr Danny W F Cheung

Article: How often does colonoscopy fail to diagnose colorectal cancer (retrospective analysis of English Hospital Episode Statistics from 2003 to 2009)?

Date: 31st August 2015

The purpose of this form is to identify all potential conflicts of interests that arise from financial relationships between any author for this article and any commercial or proprietary entity that produces healthcare-related products and/or services relevant to the content of the article. This includes any financial relationship within the last twelve months, as well as known financial relationships of authors' spouse or partner. **The lead author is responsible for submitting the disclosures of all listed authors, and must sign this form at the bottom. Additional forms may be submitted if the number of authors exceeds the space provided.**

Lead Author: Dr Danny W F Cheung

Email Address*: danny.cheung1@nhs.net

- ☒ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

Author: Felicity Evison

Email Address*: Felicity.Evison@uhb.nhs.uk

- ☒ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

Author: Mr Prashant Patel

Email Address*: Prashant.Patel@uhb.nhs.uk

- ☒ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

Author: Dr Nigel J Trudgill

Email Address*: nigel.trudgill@nhs.net

- ☒ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

* We will use email addresses only for questions related to this article 1

** **Type of relationship may include:** full-time or part-time employee, independent contractor, consultant, research or other grant recipient, paid speaker or teacher, membership on advisory committee or review panels, ownership interest (product royalty/licensing fees, owning stocks, shares, etc.), relationship of a spouse or partner, or any other financial relationship.



Author:

Email Address*:

- ☒ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

Author:

Email Address*:

- ☐ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

Author:

Email Address*:

- ☐ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

Author:

Email Address*:

- ☐ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

Author:

Email Address*:

- ☐ No financial relationships with a commercial entity producing health-care related products and/or services relevant to this article.

Company	Type of Relationship**	Content Area (if applicable)

- ☒ **As lead author of this article,** I attest that I have received disclosure information from all participating authors as listed above, and have submitted the information completely and accurately as it was reported to me. I understand that typing my name below serves as an electronic signature for the purposes of this form.

Type Name (Electronic Signature)

* We will use email addresses only for questions related to this article

** **Type of relationship may include:** full-time or part-time employee, independent contractor, consultant, research or other grant recipient, paid speaker or teacher, membership on advisory committee or review panels, ownership interest (product royalty/licensing fees, owning stocks, shares, etc.), relationship of a spouse or partner, or any other financial relationship.

January 12th 2016

Michael B. Wallace, MD, MPH
Editor-in-Chief
Seth Andrew Gross, MD
Associate Editor
GIE Editorial Team

Dear Drs. Wallace and Gross,

Re – GIE-D-15-01290: How often does colonoscopy fail to diagnose colorectal cancer (retrospective analysis of English Hospital Episode Statistics from 2003 to 2009)?

Thank you very much for sending your further editorial comments and requesting review and resubmission of our manuscript. We have detailed below our responses to the editorial comments and related amendments to the paper.

Editorial comments

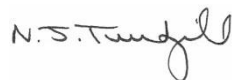
In the first paragraph of the discussion we explain the potential reasons for our reported PCCRC rate of 12.1% appearing higher than other published studies. The most important reason is that we chose to study PCCRC for five years after colonoscopy, rather than three years as some studies have done. Using a three year follow up period after colonoscopy, our PCCRC rate of 8.3% is consistent with other studies as we discuss.

Unfortunately, we are unable to provide the data requested for the period 2010 to 2014. This time period would not allow five years of follow up within the database to ascertain whether colorectal cancer developed following the colonoscopy.

However, the rate of PCCRC fell during the period we studied and we do accept the point made that there have been a number of advances in colonoscopy in recent years that should impact further on the rate of PCCRC in recent years when it is subsequently analysed. We have amended the discussion to acknowledge that the current PCCRC rate is likely to be even lower than the reported PCCRC rate in our study due to changes in colonoscopy practice. Text amended.

We hope our manuscript is now suitable for publication.

Yours sincerely,

A handwritten signature in black ink, appearing to read 'N. S. Trudgill', with a stylized flourish at the end.

Dr Nigel Trudgill

**Factors associated with colorectal cancer occurrence after colonoscopy that did not
diagnose colorectal cancer.**

¹Danny Cheung, ²Felicity Evison, ³Prashant Patel, ¹Nigel Trudgill

¹Department of Gastroenterology, Sandwell General Hospital, Lyndon, West Bromwich

²Health Informatics Department, Queen Elizabeth Hospital, Birmingham

³School of Cancer Sciences, University of Birmingham, Birmingham

Corresponding author:

Dr NJ Trudgill

Sandwell General Hospital

Lyndon

West Bromwich

B71 4HJ

Tel 0121 5073080

Fax 0121 5073265

Email nigel.trudgill@nhs.net

Word count: 3561

Abstract

Background and Aims: Up to 6% of colorectal cancers (CRC) are diagnosed within 5 years of a colonoscopy that did not diagnose CRC (post-colonoscopy colorectal cancer, PCCRC). PCCRC and associated risk factors were examined within a national hospital episode database.

Methods: A retrospective case-control study of all adult colonoscopies recorded in Hospital Episode Statistics (HES) between 2003-2009 in England. PCCRC cases underwent colonoscopy 6-60 months before diagnosis; controls had not undergone colonoscopy 6-60 months before diagnosis. Multivariate logistic regression analysis examined associations with PCCRC.

Results: 1,439,684 colonoscopies were analysed, including 67,202 CRC and 8147 (12.1%) PCCRC cases. Multivariate analysis revealed that female gender (odds ratio 1.13 (95% CI 1.08-1.19), $p<0.001$), older age (70-74 years) (1.09 (1.00-1.18), $p=0.039$), increased co-morbidity (Charlson index 5+) (1.16 (1.05-1.28), $p<0.003$) and right sided CRC (1.17 (1.11-1.23), $p<0.0001$) were associated with PCCRC. Emergency colonoscopy (0.54 (0.59-0.69), $p<0.0001$) was negatively associated with PCCRC. More PCCRC subjects developed metastases within 12 months and less underwent surgery (0.33 (0.32-0.35), $p<0.0001$) or chemotherapy (0.66 (0.62-0.69), $p<0.0001$). PCCRC rates varied twofold between providers, and was associated with medium volume providers compared with high volume (1.13 (1.01-1.27), $p=0.035$). The PCCRC rate fell from 13.8% in 2003 to 11.9% in 2009.

Conclusions: PCCRC occurred in 12.1% of CRC patients between 2003 and 2009. PCCRC was associated with female gender, older age, increased co-morbidity, right sided CRC, elective procedures and colonoscopy volume. PCCRC was associated with worse outcomes.

Introduction

Colonoscopy is the gold standard for diagnosing, screening and surveillance for CRC. In England, the setting of national standards for colonoscopy and accreditation of endoscopy units has resulted in an improvement in auditable colonoscopy standards over the last decade.[1] The same period has also coincided with an increase in 5 year survival following CRC diagnosis from 47.8% to 53.6%.[2] However, 2.6 to 6.0% of CRC patients have previously been reported to be diagnosed within 5 years of a colonoscopy which did not detect cancer. These events are termed post-colonoscopy colorectal cancer (PCCRC).[3, 4, 5] It has been proposed that PCCRC may have a different cell biology from other CRC with more aggressive and rapidly growing tumours.[6, 7] However, two recently published North American studies concluded that this did not apply to the majority of PCCRC, with around two thirds of PCCRC a result of missed lesions or incomplete polypectomy.[4, 8]

Given the improvements in colonoscopy over the past decade in England, we have examined the impact on PCCRC in a national hospital episode database and associated risk factors for these events.

Methods

Data sources

Hospital Episode Statistics (HES) is an administrative database which records information on all elective and emergency care episodes in National Health Service (NHS) hospitals in England.[9] Each care episode record includes demographic, admission, diagnoses and procedures data. Diagnoses are coded using International Classification of Diseases version 10 (ICD-10) and procedures are coded using Office of Population Censuses and Surveys Classification of Interventions and Procedures 4th revision (OPCS-4). HES is linked to Office for National Statistics (ONS) mortality records, which include date of death and causes of death recorded on death certificates. The NHS provides comprehensive healthcare coverage for the UK population, with the vast majority of colonoscopies performed in the UK by a NHS provider.[1]

Subject definitions

All subjects over the age of 18 years undergoing colonoscopy between April 2003 and March 2009 were identified from HES. Colonoscopy and CRC were defined by OPCS-4 (*appendix 1*) and ICD-10 codes (*appendix 2*) respectively. Subjects with a CRC diagnosis prior to the first episode of colonoscopy and subjects with a diagnosis of inflammatory bowel disease (IBD) were excluded from the analysis to avoid confounding through surveillance.

Recording of a CRC diagnosis in HES records may be delayed by a few weeks from the date of the diagnostic colonoscopy code.[10, 11] For the purpose of this study, the diagnosis date was therefore defined as the first colonoscopy code during the 6 months prior to the first CRC coding episode in HES or mortality records[10, 12], or the first CRC episode for those subjects who did not have a colonoscopy during this 6 month period due to being diagnosed through an alternative method, e.g. barium enema, CT colonography or flexible sigmoidoscopy. Subjects undergoing colonoscopy 6 to 60 months before subsequent CRC diagnosis were identified as PCCRC cases. These cases were further classified as PCCRC 6-12 months (colonoscopy 6 to 12 months prior to CRC diagnosis); PCCRC 12-36 months (colonoscopy 12 to 36 months prior to CRC diagnosis) and PCCRC 36-60 months

(colonoscopy 36 to 60 months prior to CRC diagnosis). For patients who had more than one colonoscopy 6 to 60 months prior to CRC diagnosis, data from the most recent colonoscopy was used for analysis. Controls were subjects who had not undergone colonoscopy in the period 6 to 60 months before CRC diagnosis. Colonoscopies from 2003 to 2009 were studied to ensure all subjects had at least 5 years of follow up within HES. The PCCRC rate was calculated from the number of PCCRC subjects divided by the sum of PCCRC subjects and controls.[13]

Validation of colonoscopy and colorectal cancer populations

To assess the validity of the HES colonoscopy population, the number of colonoscopies between 2007 and 2010 at University Hospital Birmingham (UHB) was extracted from endoscopy records (Unisoft Medical Systems, Enfield, Middlesex, UK) and compared with the number of colonoscopies recorded in HES for UHB. To assess the validity of a CRC diagnosis in HES using the study methodology, the number of HES CRC cases was compared with the number of CRC cases diagnosed in England from the National Cancer Intelligence Network (NCIN)[14] from 2002 to 2011. Finally, the rate of surgery in the HES CRC population was compared with rate of surgery in the National Bowel Cancer Audit between 2008 and 2011.[15, 16, 17]

Study variables

Subject demographics

Study variables were extracted from coding at the time of PCCRC colonoscopy in cases and diagnostic colonoscopy or first CRC episode in controls. Ethnicity was identified from HES demographic fields and grouped into White or White British, Asian or Asian British, Black or Black British, Chinese, Mixed and other ethnic groups.

Co-morbidity

The Charlson co-morbidity index was calculated using ICD-10 codes for secondary diagnoses, excluding metastatic disease, and divided into three categories: 0 (no co-morbidity), 1-4 (low co-morbidity) and 5 or greater (high co-morbidity).[18]

Socio-economic status

Deprivation was assessed using the Index of Multiple Deprivations 2007, which is an aggregate score for each English catchment area. Subjects were linked to their corresponding catchment area by postcode of residence and associations with deprivation were analysed in quintiles, with quintile 1 being the most deprived.

Colorectal cancer variables

CRC site was classified based on the first CRC coding episode into: right sided, left sided and unspecified (*appendix 3*). Coding records of initially unspecified site CRC were examined and if a more specific code had been used subsequently, this was used to determine the CRC site. Colonic polyps were identified from ICD-10 codes (*appendix 4*).

Distant metastases were identified by ICD-10 codes (*appendix 5*) up to 12 months from diagnosis date and were used as a surrogate marker of CRC stage at diagnosis, as Dukes' staging is not recorded in HES. Codes for metastases can occasionally be miscoded as a primary neoplasm (e.g. lung), and therefore primary malignancy codes were also used, provided that they were recorded in the 12 months subsequent to CRC diagnosis (*appendix 5*). Surgery and chemotherapy were identified by respective OPCS-4 codes (*appendix 6*).

Survival analysis

Survival analysis adjusted for gender, age, deprivation and co-morbidity was calculated from the CRC diagnosis date of PCCRC cases and controls using date of death from ONS. Subjects who were not diagnosed by colonoscopy were not included to avoid potential lead time bias due to the method of determining date of diagnosis from HES.

Provider variables

For the purpose of this study, all endoscopy units operating within the same NHS organisation were analysed as a single provider. Individual providers were stratified by colonoscopy volume, bowel cancer screening program (BCSP) status and the percentage of CRC diagnosed during an emergency rather than an elective episode to determine if there was an association with PCCRC. Colonoscopy volume was determined from the total number of colonoscopies performed during the study period at each provider and separated into

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

tertiles. A BCSP accredited provider had at least one endoscopy unit accredited with BCSP status by the end of the study period. The percentage of CRC diagnosed as an emergency at a provider was the ratio of CRC diagnosed during an acute (unplanned) admission divided by all CRC, including CRC diagnosed during an elective episode.

Ethics

As only pseudonymised information was used in this study, ethics approval was not necessary. HES data is available under a data sharing agreement for the purposes of service evaluation.

Statistical methodology

Statistical analysis was carried out with STATA SE v13.1 (Statacorp LP, Texas, USA). Analysis of variance and χ^2 tests were used to compare differences in continuous and categorical variables respectively. Associations with PCCRC were examined by univariate and multivariate logistic regression. A multivariate model was constructed to determine associations with PCCRC following adjusting gender, age, Charlson co-morbidity index, procedure type (emergency or elective), CRC site (left colon or right colon), metastases and procedure year. For tests of significance, p values <0.05 were considered significant. All odds ratios, 95% confidence intervals and associated p values are the result of multivariate analysis unless stated otherwise. Unadjusted Kaplan-Meier analysis and Cox proportional hazards modelling following adjustment for gender, age, deprivation and co-morbidity were used to compare survival in PCCRC cases and controls.

Results

Study cohort

Between April 2003 and March 2009, 1,439,684 colonoscopies were identified and 67,202 subjects were diagnosed with CRC during this period. Out of the 67,202 CRC subjects, there were 8,147 (12.1%) PCCRC subjects: 1796 (2.7%) PCCRC 6-12 months; 3,772 (5.6%) PCCRC 12-36 months and 2,579 (3.8%) PCCRC 36-60 months. 59,055 CRC subjects had not had a colonoscopy between 6 and 60 months prior to CRC diagnosis and served as controls. Overall, 0.66% or 1 in every 150 subjects developed PCCRC after a colonoscopy that did not diagnose CRC.

Validation of colonoscopy and colorectal cancer populations

The total number of colonoscopies carried out between 2007 and 2010 at UHB was 8708 and 8292 colonoscopies (95.2%) were coded in HES for UHB for the equivalent four year period. The CRC population was validated by comparing CRC cases recorded in HES (315,515) to CRC cases reported from 2002 to 2011 by NCIN (312,984)[14], showing a concordance of over 99%. The CRC population was further validated by comparing the 70.4% surgical rate for CRC from HES with the National Bowel Cancer Audit, which reported that 75.7% of CRC patients enrolled in the audit underwent surgery between 2008 and 2011.[15, 16, 17] All of the validation processes showed a good correlation between HES data and independent data sources, suggesting the study methodology was valid.

Subject characteristics

The characteristics of cases with PCCRC and CRC controls are shown in Table 1. PCCRC subjects (mean age 70.7 ± 11.4 years) were older than controls (mean age 70.2 ± 11.4 years)($p < 0.001$). The risk of PCCRC appeared to increase with age on univariate analysis, but only subjects aged 70 to 74 were associated with PCCRC compared with subjects under 60, following adjusting for confounding factors. PCCRC subjects were more likely to be female. Subjects with the most co-morbidities (Charlson co-morbidity index of 5 or greater) were associated with PCCRC. PCCRC was not associated with differences in ethnicity or deprivation.

Colonoscopy variables and findings

The influence of colonoscopy variables and findings on PCCRC are shown in Table 2. The majority of CRC were diagnosed during an elective colonoscopy. However, being diagnosed during an emergency colonoscopy reduced the risk of PCCRC nearly by half. There was minor increased risk of PCCRC on univariate analysis in colonoscopies carried out at the weekend compared with during the week.

PCCRC was more likely to be associated with CRC in the right colon. Colonic polyps were coded in 21.6% of the colonoscopies which did not detect CRC in the PCCRC group.

Polypectomy was coded in a further 18.9%. On univariate analysis, this was higher than both the recorded polyp rate of 9.8% (2.52 (95% CI 2.39-2.65), $p<0.0001$) and polypectomy rate of 11.3% (1.82 (95% CI 1.72-1.92), $p<0.0001$) from all colonoscopies during the study period. Furthermore, the polyp and polypectomy rates were both higher in the PCCRC 6-12 months group on univariate analysis, than in the PCCRC 12-36 months ($p<0.0001$) and PCCRC 36-60 months ($p<0.0001$) groups.

Colorectal outcomes and survival

The prevalence of metastatic disease within 12 months of CRC diagnosis in PCCRC cases and controls are shown in Table 3. PCCRC cases were up to twice as likely to be diagnosed with lung, peritoneal and bone metastases within 12 months of CRC diagnosis. However, lymph node metastases were more prevalent in controls than PCCRC cases, suggesting coding bias related to the increased rate of surgery in control subjects described later.

On univariate analysis, PCCRC cases were less likely to undergo surgery compared with controls (0.33 (95% CI 0.32-0.35), $p<0.0001$) or chemotherapy (0.66 (95% CI 0.62-0.69), $p<0.0001$). Overall survival was also worse in PCCRC subjects compared with controls, with a median survival of 5.8 years in controls compared with 2.1 years in the PCCRC 6-12 months group, 2.0 years in the PCCRC 12-36 months group and 3.5 years in the PCCRC 36-60 months group (figure 1). Following adjusting for age, gender, co-morbidity and deprivation, survival outcomes remained worse for PCCRC subjects with a hazard ratio of 1.17 (95% CI 1.10-1.24)($p<0.0001$), 1.26 (95% CI 1.20-1.31)($p<0.0001$) and 1.20 (95% CI 1.13-1.27)($p<0.0001$)

for the PCCRC 6-12 months, PCCRC 12-36 months and PCCRC 36-60 months respectively when compared with controls.

Individual provider variables

The influence of provider variables on PCCRC are shown in Table 4. There was a more than twofold variation in PCCRC rates between individual providers in England during the study period (figure 2). On univariate analysis, medium colonoscopy volume providers and low volume providers were both more likely to be associated with PCCRC than high volume providers. Following adjusting for other variables in the multivariate model an association with medium volume providers remained. BCSP accreditation status and the percentage of CRC diagnosed as an emergency were not associated with an increased risk of PCCRC.

PCCRC rates over time

The number of colonoscopies recorded in HES has increased by almost two fold over the study period. Despite the increase in colonoscopy numbers performed year on year, the annual rate of PCCRC has steadily fallen over the study period ($p < 0.0001$)(figure 3). The annual PCCRC rate decreased from 13.8% in 2003-2004 to 11.9% by the end of study period in 2008-2009 with the reduction seen mainly in the PCCRC 6-12 months and PCCRC 12-36 months groups.

Discussion

The overall PCCRC rate of 12.1% in 67202 subjects in England between 2003 and 2009 appears higher than previously published figures. However, some previous studies have calculated the PCCRC rate by only including CRC subjects with a colonoscopy up to 36 months prior to diagnosis and the comparable figure from the present study is 8.3%. A Canadian study of 14,064 CRC subjects reported a PCCRC rate of 9.0% between 2000 and 2005.[12] Using the Surveillance, Epidemiology, and End Results (SEER)-Medicare database in the USA, a PCCRC rate of 7.2% was reported between 1994 to 2005 from a study of 57,839 CRC subjects.[19] A further population based study from Utah, USA with 2659 CRC subjects between 1995 and 2009 described a PCCRC rate of 6% when subjects with a colonoscopy up to 60 months prior to CRC diagnosis were included.[4] In Europe, two recent studies have reported much lower PCCRC rates. A Danish population based study between 2000 to 2009 included 37,044 CRC subjects and concluded that only 2.7% of CRC subjects have had a colonoscopy that failed to diagnose CRC 1 to 5 years prior to diagnosis.[5] A second study from the Netherlands analysed 5107 CRC subjects between 2001 to 2010 from three providers and found a PCCRC rate of only 2.9% for subjects with a colonoscopy up to 60 months prior to CRC diagnosis.[20] In addition to potential variations in subject and colonoscopy factors between the difference studies, the wide range of reported PCCRC rates are likely to be contributed to by methodological differences.[13]

In the present study, PCCRC was associated with older subjects, female gender, an increased number of co-morbidities and right-sided CRC, which is in keeping with findings from other studies of PCCRC. [3, 12, 19, 21] The association between increasing age and PCCRC was less marked on multivariate analysis and this may relate to confounding from increasing co-morbidity in the elderly. Elderly patients are more likely to have inadequate bowel preparation, thus reducing mucosal visualisation and detection of polyps and early CRC.[22, 23] Female patients are more likely to have had previous abdominal and pelvic surgery, which may increase the technical difficulty of colonoscopy and impair patient tolerance, reducing the caecal intubation rate.[24] In addition to factors that have an adverse effect on caecal intubation rate, right sided CRC are more likely to arise from flat, non-polypoid

adenomatous lesions[20, 25] that poor bowel preparation may make difficult to detect. This will contribute to the association of right sided CRC with PCCRC.

Over a fifth of PCCRC subjects had colonic polyps or polypectomy coded during the most recent colonoscopy prior to CRC diagnosis. This is higher than the average polypectomy rate in all colonoscopy procedures during the same period. Furthermore, polyp and polypectomy coding rates were highest in the PCCRC 12-36 months group. Prior polypectomy has been reported to double the risk of PCCRC[19], with up to 19% of CRC occurring in the same anatomic segment as a previously resected adenoma.[8] Paradoxically, colonoscopists with higher polypectomy rates have been reported to be associated with a lower risk of PCCRC[12, 19], presumably as they detect more polyps and remove them more completely than other colonoscopists. Incomplete polypectomy, or inadequate biopsy sampling of polyps, is therefore a key modifiable risk factor for PCCRC and ensuring adequate follow up and assessment following polypectomy may reduce PCCRC rates.

PCCRC subjects appeared to have worse outcomes in terms of both treatment following diagnosis (surgery and chemotherapy) and overall survival. Previous studies have reported no survival difference between PCCRC subjects and controls[5, 21] with one recent study even reporting a survival benefit in the PCCRC subjects, which was likely to be due to earlier CRC stage at diagnosis in the PCCRC subjects.[4] In the current study, PCCRC subjects were older, had greater co-morbidities and were more likely to present with distant metastases within 12 months of diagnosis compared with controls. All these factors contributed to the reduced rates of curative surgery or palliative chemotherapy for PCCRC subjects and will have contributed to worse survival. Adjusting the survival analyses for differences in ages, gender, co-morbidity and deprivation still revealed worse survival for PCCRC subjects and, at least in England, PCCRC is clearly associated with worse survival. Survival in PCCRC subjects would have been potentially better if earlier opportunities to diagnose their CRC had been taken.

Previous studies have reported that PCCRC was not associated with endoscopist procedure volume[12] and that higher colonoscopy volumes may even be positively associated with PCCRC surprisingly.[19] In the current study, there was a large variation in PCCRC rates

1 between individual providers across England but PCCRC appeared to be associated with
2 lower colonoscopy volume providers. This result should be interpreted with caution. We did
3 not have access to colonoscopy quality indicators such as caecal intubation and adenoma
4 detection rates that are likely to be potentially more important factors in PCCRC incidence.
5
6
7
8
9

10 Colonoscopy undertaken during an emergency admission covered 10% of procedures
11 examined and was associated with a lower risk of PCCRC at 9% compared with 14% for
12 elective procedures. Patients presenting as an emergency may have more advanced
13 colorectal cancer and therefore a lower chance of PCCRC.
14
15
16
17
18

19 The annual PCCRC rate in England has fallen steadily over the study period from 13.8% to
20 11.9%, at least partly due to improving colonoscopy standards over the corresponding time
21 period. In 2003, a multi-regional audit in England including 9223 colonoscopies reported
22 that mean caecal intubation rate was only 76.9%.[26] A subsequent national audit in 2011
23 of 20085 colonoscopies found that the caecal intubation rate had improved to 92.3%.[1] The
24 PCCRC rate is likely to continue to improve in recent years given changes in colonoscopy
25 practice, including the recognition of the importance of minimum withdrawal times [27],
26 bowel preparation improvements[28] and better endoscopic recognition of sessile serrated
27 polyps[25], subsequent to the study period.
28
29
30
31
32
33
34
35
36
37
38

39 The use of a national hospital dataset enabled us to undertake one of the largest PCCRC
40 studies to date, including the vast majority of colonoscopies performed during a period of
41 rising colonoscopy standards. The quality of diagnostic and procedural coding in HES has
42 been previously investigated and there was a high concordance when compared with
43 independent national data sources.[1, 10, 29] However, we did not have the opportunity to
44 link our HES dataset directly to cancer registry data due to restrictions under which the data
45 is held and therefore, in order to validate the methodology chosen, colonoscopy and CRC
46 populations were compared with national cancer databases and a local data sample and
47 revealed a good correlation. The completeness and accuracy of coding in HES is still a
48 potential source of concern. For example, the diagnosis date may not be recorded
49 accurately in HES due to the need for histological confirmation before CRC coding and
50 therefore a colonoscopy within 6 months of CRC coding had to be considered the diagnostic
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

1 procedure. There are also limitations in HES concerning coding of colonoscopy procedures,
2 polyps, polypectomy, presence of metastases, surgery and chemotherapy and the figures
3 included may be an over or under estimate, though this is likely to affect PCCRC cases and
4 controls equally. A further limitation is that key procedure information such as the bowel
5 preparation quality, sedation doses, colonoscopist grade and specialty, extent of
6 examination, completeness of polypectomy and number of biopsies taken are not recorded
7 in HES and all may influence the PCCRC risk. Furthermore, due to the HES coding hierarchy,
8 indication, presence of diverticular disease and history of abdominal or pelvic surgery may
9 not be coded, partly due to under reporting by colonoscopists when significant pathology or
10 CRC are found and again each may be important risk factors for PCCRC. As HES does not
11 record polyp histology or the International Classification of Diseases for Oncology (ICD-O)
12 codes, the lack of data on polyp and CRC histology and Duke's staging further limits analysis
13 of potential causes of PCCRC (de novo CRC, incomplete adenoma resection, missed lesion or
14 biopsy failed to detect CRC) and survival in PCCRC subjects.
15
16
17
18
19
20
21
22
23
24
25
26
27
28

29 In conclusion, the PCCRC rate was 12.1% in England between 2003 and 2009. PCCRC was
30 associated with older age, female gender, increasing co-morbidity, procedure related
31 factors (elective procedures and right sided CRC) and provider colonoscopy volume. Despite
32 the encouraging fall in annual PCCRC rate over the study period, PCCRC rate should be a
33 routinely measured endoscopy unit colonoscopy quality marker and potentially avoidable
34 risk factors for PCCRC addressed.
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

Table 1. The characteristics of post-colonoscopy colorectal cancer cases and controls

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	p value	Odds ratio	95% CI	p value
Total subjects (number)						Univariate			Multivariate		
	1796 (2.7)	3772 (5.6)	2579 (3.8)	8147 (12.1)	59055 (87.9)	-	-	-	-	-	-
Mean age \pmSD (years)											
	71.5 \pm 11.4	70.9 \pm 11.7	69.8 \pm 10.8	70.7 \pm 11.4	70.2 \pm 11.4	-	-	<0.001			
Age group (number (%))											
Under 60	263 (3.2)	598 (7.3)	415 (5.1)	1276 (15.7)	9849 (16.7)	Ref					
60-64	167 (2.0)	367 (4.5)	288 (3.5)	822 (10.1)	6749 (11.4)	0.94	0.86-1.03	0.1928	0.95	0.86-1.04	0.277
65-69	217 (2.7)	531 (6.5)	435 (5.3)	1183 (14.5)	8810 (14.9)	1.04	0.95-1.13	0.4044	1.03	0.94-1.12	0.537
70-74	344 (4.2)	648 (8.0)	488 (6.0)	1480 (18.2)	10229 (17.3)	1.12	1.03-1.21	0.0067	1.09	1.00-1.18	0.039
75-79	359 (4.4)	678 (8.3)	499 (6.1)	1536 (18.9)	10698 (18.1)	1.11	1.02-1.20	0.0109	1.07	0.98-1.16	0.159
80+	446 (5.5)	950 (11.7)	454 (5.6)	1850 (22.7)	12720 (21.5)	1.12	1.04-1.21	0.0029	1.08	1.00-1.17	0.065
Gender (number (%))											
Male	974 (12.0)	1974 (24.2)	1340 (16.4)	4288 (52.6)	33057 (56.0)	Ref	-	-	Ref	-	-
Female	822 (10.1)	1798 (22.1)	1239 (15.2)	3859 (47.4)	25998 (44.0)	1.14	1.09-1.20	<0.0001	1.13	1.08-1.19	<0.001
Charlson co-morbidity index (number (%))											
0	1514 (18.6)	3210 (39.4)	2235 (27.4)	6959 (85.4)	50663 (85.8)	Ref	-	-	Ref	-	-
1-4	154 (1.9)	298 (3.7)	210 (2.6)	662 (8.1)	4764 (8.1)	1.01	0.93-1.10	0.7896	1.06	0.97-1.16	0.195
5+	128 (1.6)	264 (3.2)	134 (1.6)	526 (6.5)	3628 (6.1)	1.06	0.96-1.16	0.2641	1.16	1.05-1.28	0.003
Deprivation quintile (number (%))											
1 (most)	329 (4.0)	637 (7.8)	393 (4.8)	1359 (16.7)	10015 (17.0)	Ref	-	-	-	-	-
2	365 (4.5)	740 (9.1)	499 (6.1)	1604 (19.7)	11258 (19.1)	1.05	0.97-1.13	0.2153	-	-	-
3	333 (4.1)	782 (9.6)	551 (6.8)	1666 (20.4)	12399 (21.0)	0.99	0.91-1.07	0.8002	-	-	-
4	387 (4.8)	784 (9.6)	568 (7.0)	1739 (21.3)	12642 (21.4)	1.01	0.94-1.09	0.7242	-	-	-
5 (least)	381 (4.7)	823 (10.1)	566 (6.9)	1770 (21.7)	12620 (21.4)	1.03	0.96-1.11	0.3905	-	-	-
Ethnicity (number (%))											
Caucasian	1656 (20.3)	3536 (43.4)	2467 (30.3)	7659 (94.0)	54512 (92.3)	Ref	-	-	-	-	-
Asian	21 (0.3)	55 (0.7)	36 (0.4)	112 (1.4)	788 (1.3)	1.01	0.83-1.23	0.9097	-	-	-

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49

Afro-Caribbean	25 (0.3)	53 (0.7)	27 (0.3)	105 (1.3)	823 (1.4)	0.91	0.74-1.11	0.3553	-	-	-
Chinese	0	0	0	12 (0.1)	118 (0.2)	0.72	0.40-1.30	0.2865	-	-	-
Mixed	0	0	0	18 (0.2)	160 (0.3)	0.80	0.49-1.30	0.3719	-	-	-
Others	12 (0.1)	21 (0.3)	21 (0.3)	54 (0.7)	341 (0.6)	1.13	0.85-1.50	0.4156	-	-	-
Unknown	74 (0.9)	95 (1.2)	18 (0.2)	187 (2.3)	2313 (3.9)	0.58	0.49-0.67	<0.0001	-	-	-

Odds ratios with 95% confidence intervals and p values for PCCRC (all) compared with controls
PCCRC – post-colonoscopy colorectal cancer

Table 2. The colonoscopy characteristics and findings of post-colonoscopy colorectal cancer cases and controls

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	p value	Odds ratio	95% CI	p value
Procedure day (number (%))						Univariate			Multivariate		
Weekday	1736 (21.3)	3628 (44.5)	2486 (30.5)	7850 (96.4)	57249 (96.9)	Ref	-	-	-	-	-
Weekend	60 (0.7)	144 (1.8)	93 (1.1)	297 (3.6)	1806 (3.1)	1.19	1.06-1.36	0.0044	-	-	-
Procedure type (number (%))											
Elective	1622 (19.9)	3473 (42.6)	2455 (30.1)	7550 (92.7)	52605 (89.1)	Ref	-	-	Ref	-	-
Emergency	174 (2.1)	299 (3.7)	124 (1.5)	597 (7.3)	6450 (10.9)	0.64	0.59-0.70	<0.0001	0.54	0.59-0.69	<0.0001
Colorectal cancer location (number (%))											
Left sided	897 (11.0)	1754 (21.5)	1260 (15.5)	3911 (48.0)	34703 (58.8)	Ref	-	-	Ref	-	-
Right sided	535 (6.6)	1242 (15.2)	919 (11.3)	2696 (33.1)	20751 (35.1)	1.15	1.09-1.21	<0.0001	1.17	1.11-1.23	<0.0001
Unknown/overlapping sites	364 (4.5)	776 (9.5)	400 (4.9)	1540 (18.9)	3601 (6.1)	3.79	3.54-4.06	<0.0001	3.72	3.46-3.99	<0.0001
Polyp/ polypectomy coded (number (%))											
Polyp coded	491 (6.0)	742 (9.1)	523 (6.4)	1756 (21.6)	141799* (9.8)	2.52⁺	2.39-2.65⁺	<0.0001⁺	-	-	-
No polyp coded	1305 (16.0)	3030 (37.2)	2056 (25.2)	6391 (78.4)	1300714* (90.2)	Ref	-	-	-	-	-
Polypectomy coded	348 (4.3)	669 (8.2)	523 (6.4)	1540 (18.9)	162364* (11.3)	1.82⁺	1.72-1.92⁺	<0.0001⁺	-	-	-
No polypectomy coded	1448 (17.8)	3103 (38.1)	2056 (25.2)	6607 (81.1)	1280150* (89.7)	Ref	-	-	-	-	-

Odds ratios with 95% confidence intervals and p values for all PCCRC compared with controls

PCCRC – post-colonoscopy colorectal cancer

* From all colonoscopies

+ Univariate analysis comparing all PCCRC with all colonoscopies during study period.

Table 3. The prevalence of metastases within 12 months of colorectal cancer diagnosis in post-colonoscopy colorectal cancer cases and controls

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	p value	Odds ratio	95% CI	p value
Subjects with metastases within 12 months of diagnosis (number (%))						Univariate			Multivariate		
Liver metastases	276 (3.4)	619 (7.6)	365 (4.5)	1260 (15.5)	8545 (14.5)	1.08	1.01-1.15	0.017	0.97	0.91-1.05	0.486
Lung metastases	154 (1.9)	345 (4.2)	182 (2.2)	681 (8.4)	3104 (5.3)	1.64	1.51-1.79	<0.0001	1.61	1.46-1.77	<0.0001
Peritoneal metastases	75 (0.9)	166 (2.0)	102 (1.3)	343 (4.2)	1903 (3.2)	1.32	1.17-1.48	<0.0001	1.27	1.12-1.44	<0.0001
Bone metastases	45 (0.6)	106 (1.3)	78 (1.0)	229 (2.8)	678 (1.1)	2.49	2.14-2.90	<0.0001	2.21	1.88-2.60	<0.0001
Lymph node metastases	136 (1.7)	282 (3.5)	231 (2.8)	649 (8.0)	6459 (10.9)	0.70	0.65-0.76	<0.0001	0.75	0.69-0.82	<0.0001
Treatment outcome following diagnosis (number (%))											
Surgery	791 (9.7)	1661 (20.4)	1337 (16.4)	3789 (46.5)	42790 (72.5)	0.33	0.32-0.35	<0.0001	-	-	-
Chemotherapy	422 (5.2)	911 (11.2)	594 (7.3)	1927 (23.7)	18908 (32.0)	0.66	0.62-0.69	<0.0001	-	-	-

Odds ratios with 95% confidence intervals and p values for PCCRC (all) compared with controls

PCCRC – post-colonoscopy colorectal cancer

Table 4. The influence of provider variables on post-colonoscopy colorectal cancer

	PCCRC 6-12 months	PCCRC 12-36 months	PCCRC 36-60 months	All PCCRC	Controls	Odds ratio	95% CI	p value	Odds ratio	95% CI	p value
Colonoscopy volume by NHS provider (number (%))						Univariate			Multivariate		
High volume providers (>1680 pa)	955 (11.7)	1993 (24.5)	1415 (17.4)	4363 (53.6)	33353 (56.5)	Ref	-	-	Ref	-	-
Medium volume providers	704 (8.6)	1486 (18.2)	994 (12.2)	3184 (39.1)	21942 (37.2)	1.11	1.06-1.16	<0.0001	1.13	1.01-1.27	0.035
Low volume providers (<747 pa)	137 (1.7)	293 (3.6)	170 (2.1)	600 (7.4)	3760 (6.4)	1.22	1.11-1.34	<0.0001	1.05	0.98-1.12	0.161
BCSP status (number (%))											
BCSP provider	959 (11.8)	2064 (25.3)	1396 (17.1)	4419 (54.2)	31780 (53.8)	Ref	-	-	-	-	-
Non-BCSP provider	837 (10.3)	1708 (21.0)	1183 (14.5)	3728 (45.8)	27275 (46.2)	0.98	0.94-1.03	0.4690	0.96	0.90-1.03	0.255
Percentage of CRC diagnosed during an emergency admission by NHS provider (number (%))											
Low percentage providers (<27.3%)	408 (5.0)	848 (10.4)	629 (7.7)	1885 (23.1)	14270 (24.2)	0.91	0.84-0.98	0.0115	0.96	0.87- 1.06	0.443
Medium percentage providers	1068 (13.1)	2273 (27.9)	1530 (18.8)	4871 (59.8)	35211 (59.6)	0.95	0.89-1.01	0.1299	0.96	0.85-1.09	0.531
High percentage providers (>33.9%)	320 (3.9)	651 (8.0)	420 (5.2)	1391 (17.1)	9572 (16.2)	Ref	-	-	Ref	-	-

Odds ratios with 95% confidence intervals and p values for PCCRC (all) compared with controls

PCCRC – post-colonoscopy colorectal cancer

BCSP – Bowel Cancer Screening Program

Figure 1. Post-colonoscopy colorectal cancer rates by individual provider in England between 2003 and 2009.

Figure 2. Unadjusted survival following colorectal cancer diagnosis in post-colonoscopy colorectal cancer cases and control subjects.

Figure 3 Post-colonoscopy colorectal cancer rates and colonoscopy volume in England by year.

Appendix 1 - OPCS-4 codes for colonoscopy

H20.1 Snare polypectomy

H20.6 Polypectomy with colonoscopy

H22.1 Diagnostic fibreoptic endoscopic examination of colon and biopsy of lesion of colon

H22.8 Other specified diagnostic endoscopic examination of colon

H22.9 Unspecified diagnostic endoscopic examination of colon

Appendix 2 - ICD-10 codes for colorectal cancers

C18 Malignant neoplasm of colon - excluding C18.1 (malignant neoplasm of appendix)

C19 Malignant neoplasm of rectosigmoid junction

C20 Malignant neoplasm of rectum

Appendix 3 – ICD-10 codes for colorectal cancer (CRC) sites

Right sided CRC

C18.0 Caecum, Ileocaecal valve

C18.2 Ascending colon

C18.3 Hepatic flexure

C18.4 Transverse colon

Left sided CRC

C18.5 Splenic flexure

C18.6 Descending colon

C18.7 Sigmoid colon

C19 Rectosigmoid junction

C20 Rectum

Unspecified CRC location

C18.8 Overlapping lesion of colon

C18.9 Colon, unspecified

Appendix 4 - ICD-10 codes for colorectal polyps

D12.0 Caecal polyp(s)

1 D12.2 Ascending colon polyp(s)

2 D12.3 Transverse colon, hepatic flexure, splenic flexure polyp(s)

3 D12.4 Descending colon polyp(s)

4 D12.5 Sigmoid colon polyp(s)

5 D12.6 Colon, site unspecified polyp(s)

6 D12.7 Rectosigmoid junction polyp(s)

7 D12.8 Rectal polyp(s)

8
9
10
11
12
13
14
15 **Appendix 5 - ICD-10 codes for metastases**

16 C77.1 Intrathoracic lymph nodes

17 C77.2 Intra-abdominal lymph nodes

18 C77.4 Inguinal and lower limb lymph nodes

19 C77.5 Intrapelvic lymph nodes

20 C78.0 Secondary malignant neoplasm of lung

21 C78.6 Secondary malignant neoplasm of retroperitoneum and peritoneum

22 C78.7 Secondary malignant neoplasm of liver and intrahepatic bile duct

23 C79.5 Secondary malignant neoplasm of bone and bone marrow

24 C34 Malignant neoplasm of bronchus and lung

25 C48 Malignant neoplasm of retroperitoneum and peritoneum

26 C22 Malignant neoplasm of liver

27 C40-C41 Malignant neoplasms of bone and articular cartilage

28
29
30
31
32
33
34
35
36
37
38
39
40
41
42 **Appendix 6- OPCS-4 codes for surgical procedures**

43 H04 Total excision of colon and rectum

44 H05 Total excision of colon

45 H06 Extended excision of right hemicolon

46 H07 Other excision of right hemicolon

47 H08 Excision of transverse colon

48 H09 Excision of left hemicolon

49 H10 Excision of sigmoid colon

50 H11 Other excision of colon

51 H29 Subtotal excision of colon

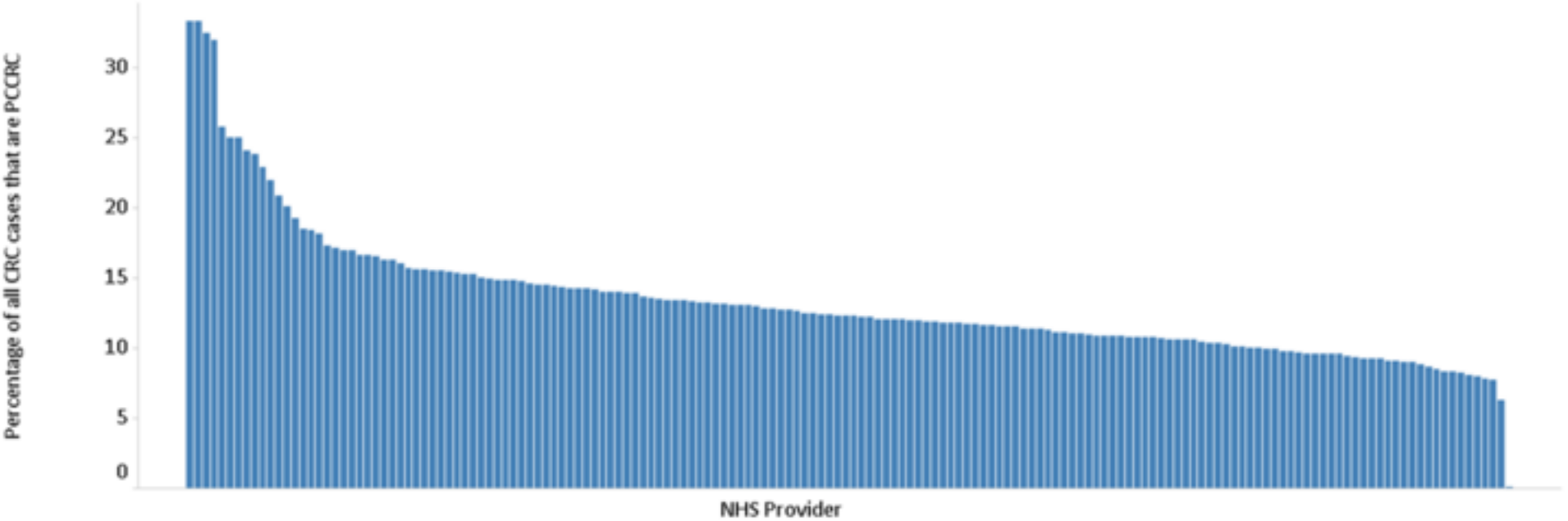
1 H33 Excision of rectum
2 H40 Operations on rectum through anal sphincter
3
4 H122 Excision of lesion of colon NEC
5
6 H123 Destruction of lesion of colon NEC
7
8 H128 Other specified extirpation of lesion of colon
9
10 H129 Unspecified extirpation of lesion of colon
11
12 H341 Open excision of lesion of rectum
13
14 H345 Open destruction of lesion of rectum
15
16 H348 Other specified open extirpation of lesion of rectum
17
18 H349 Unspecified open extirpation of lesion of rectum
19
20 H402 Trans-sphincteric excision of lesion of rectum
21
22 H403 Trans-sphincteric destruction of lesion of rectum
23
24 OPCS-4 codes for chemotherapy
25
26 X70 Procurement of drugs for chemotherapy for neoplasm in Bands 1-5
27
28 X71 Procurement of drugs for chemotherapy for neoplasm in Bands 6-10
29
30 X72 Delivery of Chemotherapy for neoplasm
31
32 X73 Delivery of oral chemotherapy for neoplasm
33
34 X352 Intravenous chemotherapy
35
36 X384 Subcutaneous chemotherapy
37
38 X373 Intramuscular chemotherapy
39
40 Z082 Follow up examination after chemotherapy for malignant neoplasm
41
42 Z511 Chemotherapy session for neoplasm
43
44 Z542 Convalescence following chemotherapy
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

References

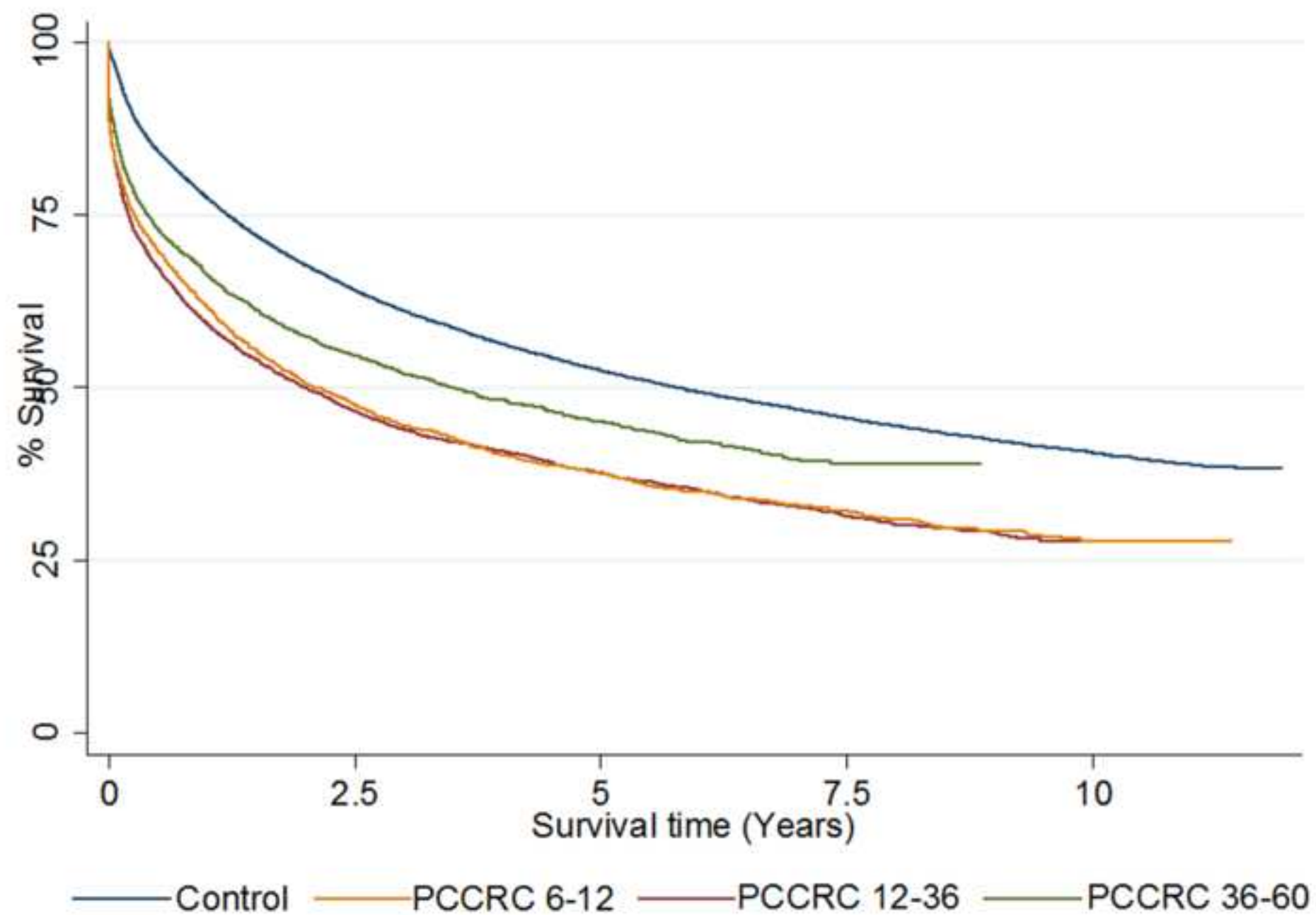
- 1 Gavin DR, Valori RM, Anderson JT, Donnelly MT, Williams JG, Swarbrick ET. The national
colonoscopy audit: a nationwide assessment of the quality and safety of colonoscopy in the UK. *Gut*
2013;**62**:242-9.
- 2 Coleman MP, Forman D, Bryant H, Butler J, Rachet B, Maringe C, *et al.* Cancer survival in
Australia, Canada, Denmark, Norway, Sweden, and the UK, 1995-2007 (the International Cancer
Benchmarking Partnership): an analysis of population-based cancer registry data. *Lancet*
2011;**377**:127-38.
- 3 Bressler B, Paszat LF, Chen Z, Rothwell DM, Vinden C, Rabeneck L. Rates of new or missed
colorectal cancers after colonoscopy and their risk factors: a population-based analysis.
Gastroenterology 2007;**132**:96-102.
- 4 Samadder NJ, Curtin K, Tuohy TM, Pappas L, Boucher K, Provenzale D, *et al.* Characteristics
of missed or interval colorectal cancer and patient survival: a population-based study.
Gastroenterology 2014;**146**:950-60.
- 5 Erichsen R, Baron JA, Stoffel EM, Laurberg S, Sandler RS, Sorensen HT. Characteristics and
survival of interval and sporadic colorectal cancer patients: a nationwide population-based cohort
study. *The American journal of gastroenterology* 2013;**108**:1332-40.
- 6 Arain MA, Sawhney M, Sheikh S, Anway R, Thyagarajan B, Bond JH, *et al.* CIMP status of
interval colon cancers: another piece to the puzzle. *The American journal of gastroenterology*
2010;**105**:1189-95.
- 7 Sawhney MS, Farrar WD, Gudiseva S, Nelson DB, Lederle FA, Rector TS, *et al.* Microsatellite
instability in interval colon cancers. *Gastroenterology* 2006;**131**:1700-5.
- 8 Robertson DJ, Lieberman DA, Winawer SJ, Ahnen DJ, Baron JA, Schatzkin A, *et al.* Colorectal
cancers soon after colonoscopy: a pooled multicohort analysis. *Gut* 2014;**63**:949-56.
- 9 Health & Social Care Information Centre. *Hospital Episode Statistics*; www.hscic.gov.uk/hes
[Accessed 29 December 2013].
- 10 Shaihi M, Thompson E, Kapoor N, Powell G, Sturges RP, Stern N, *et al.* Variation in
gastroscopy rate in English general practice and outcome for oesophagogastric cancer: retrospective
analysis of Hospital Episode Statistics. *Gut* 2014;**63**:250-61.
- 11 Weller D, Vedsted P, Rubin G, Walter FM, Emery J, Scott S, *et al.* The Aarhus statement:
improving design and reporting of studies on early cancer diagnosis. *British journal of cancer*
2012;**106**:1262-7.
- 12 Baxter NN, Sutradhar R, Forbes SS, Paszat LF, Saskin R, Rabeneck L. Analysis of administrative
data finds endoscopist quality measures associated with postcolonoscopy colorectal cancer.
Gastroenterology 2011;**140**:65-72.
- 13 Morris EJ, Rutter MD, Finan PJ, Thomas JD, Valori R. Post-colonoscopy colorectal cancer
(PCCRC) rates vary considerably depending on the method used to calculate them: a retrospective
observational population-based study of PCCRC in the English National Health Service. *Gut*
2015;**64**:1248-56.
- 14 National Cancer Intelligence Network. UK Cancer e-Atlas by cancer networks.
- 15 Health & Social Care Information Centre. *National Bowel Cancer Audit 2009*;
www.hscic.gov.uk/catalogue/PUB02587/nati-clin-audi-supp-prog-bowe-canc-2009-rep2.pdf
[Accessed 2 June 2014]. 2009.
- 16 Health & Social Care Information Centre. *National Bowel Cancer Audit 2010*;
www.hscic.gov.uk/catalogue/PUB02586/nati-clin-audi-supp-prog-bowe-canc-2010-rep.pdf
[Accessed 2 June 2014]. 2010.
- 17 Health & Social Care Information Centre. *National Bowel Cancer Audit 2011*;
www.hscic.gov.uk/catalogue/PUB02576/nati-clin-audi-supp-prog-bowe-canc-2011-rep1.pdf
[Accessed 2 June 2014]. 2011.

- 18 Charlson ME, Pompei P, Ales KL, MacKenzie CR. A new method of classifying prognostic comorbidity in longitudinal studies: development and validation. *Journal of chronic diseases* 1987;**40**:373-83.
- 19 Cooper GS, Xu F, Barnholtz Sloan JS, Schluchter MD, Koroukian SM. Prevalence and predictors of interval colorectal cancers in medicare beneficiaries. *Cancer* 2012;**118**:3044-52.
- 20 le Clercq CM, Bouwens MW, Rondagh EJ, Bakker CM, Keulen ET, de Ridder RJ, *et al*. Postcolonoscopy colorectal cancers are preventable: a population-based study. *Gut* 2014;**63**:957-63.
- 21 Singh H, Nugent Z, Demers AA, Bernstein CN. Rate and predictors of early/missed colorectal cancers after colonoscopy in Manitoba: a population-based study. *The American journal of gastroenterology* 2010;**105**:2588-96.
- 22 Romero RV, Mahadeva S. Factors influencing quality of bowel preparation for colonoscopy. *World journal of gastrointestinal endoscopy* 2013;**5**:39-46.
- 23 Hong SN, Sung IK, Kim JH, Choe WH, Kim BK, Ko SY, *et al*. The Effect of the Bowel Preparation Status on the Risk of Missing Polyp and Adenoma during Screening Colonoscopy: A Tandem Colonoscopic Study. *Clinical endoscopy* 2012;**45**:404-11.
- 24 Shah HA, Paszat LF, Saskin R, Stukel TA, Rabeneck L. Factors associated with incomplete colonoscopy: a population-based study. *Gastroenterology* 2007;**132**:2297-303.
- 25 Tadros M, Anderson JC. Serrated polyps: clinical implications and future directions. *Current gastroenterology reports* 2013;**15**:342.
- 26 Bowles CJ, Leicester R, Romaya C, Swarbrick E, Williams CB, Epstein O. A prospective study of colonoscopy practice in the UK today: are we adequately prepared for national colorectal cancer screening tomorrow? *Gut* 2004;**53**:277-83.
- 27 Simmons DT, Harewood GC, Baron TH, Petersen BT, Wang KK, Boyd-Enders F, *et al*. Impact of endoscopist withdrawal speed on polyp yield: implications for optimal colonoscopy withdrawal time. *Alimentary pharmacology & therapeutics* 2006;**24**:965-71.
- 28 Aoun E, Abdul-Baki H, Azar C, Mourad F, Barada K, Berro Z, *et al*. A randomized single-blind trial of split-dose PEG-electrolyte solution without dietary restriction compared with whole dose PEG-electrolyte solution with dietary restriction for colonoscopy preparation. *Gastrointestinal endoscopy* 2005;**62**:213-8.
- 29 Moller H, Richards S, Hanchett N, Riaz SP, Luchtenborg M, Holmberg L, *et al*. Completeness of case ascertainment and survival time error in English cancer registries: impact on 1-year survival estimates. *British journal of cancer* 2011;**105**:170-6.

Figure(1) (must be TIF or EPS files)
[Click here to download high resolution image](#)



Figure(2) (must be TIF or EPS files)
[Click here to download high resolution image](#)



Figure(3) (must be TIF or EPS files)
[Click here to download high resolution image](#)

